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(54) Title: OSTERTAGIA VACCINE

(57) Abstract: The present invention relates to nucleic acid sequences encoding *Ostertagia ostertagi* proteins and to parts of such nucleic acid sequences that encode an immunogenic fragment of such proteins, and to DNA fragments, recombinant DNA molecules, live recombinant carriers and host cells comprising such nucleic acid sequences or such parts thereof. The invention also relates to *Ostertagia ostertagi* proteins and immunogenic parts thereof encoded by such sequences. Furthermore, the present invention relates to vaccines comprising such nucleic acid sequences and parts thereof, DNA fragments, recombinant DNA molecules, live recombinant carriers and host cells comprising such nucleic acid sequences or such parts thereof, proteins or immunogenic parts thereof and antibodies against such proteins or immunogenic parts thereof. Also, the invention relates to the use of said proteins in vaccines and for the manufacture of vaccines. Moreover, the invention relates to the use of aid nucleic acid sequences, proteins or antibodies for diagnostic or vaccination purposes. Finally the invention relates to diagnostic kits comprising such nucleic acids, proteins or antibodies against such proteins.

Ostertagia vaccine

The present invention relates to nucleic acid sequences encoding *Ostertagia ostertagi* proteins, to parts of such nucleic acid sequences that encode an immunogenic fragment 5 of such proteins, to DNA fragments, recombinant DNA molecules, live recombinant carriers and host cells comprising such nucleic acid sequences or such parts thereof. The invention also relates to *Ostertagia ostertagi* proteins and immunogenic parts thereof encoded by such sequences. Furthermore, the present invention relates to vaccines comprising such nucleic acid sequences and parts thereof, DNA fragments, recombinant 10 DNA molecules, live recombinant carriers and host cells comprising such nucleic acid sequences or such parts thereof, proteins or immunogenic parts thereof and antibodies against such proteins or immunogenic parts thereof. Also, the invention relates to the use of said proteins in vaccines and for the manufacture of vaccines. Moreover, the invention relates to the use of said nucleic acid sequences, proteins or antibodies for diagnostic or 15 vaccination purposes. Finally the invention relates to diagnostic kits comprising such nucleic acids, proteins or antibodies against such proteins.

There are about 82 million cattle in the EU and about 97 million in the USA most of which are exposed to infection with gastro-intestinal nematodes at grazing, with resultant, often 20 substantial, impaired production efficiency. The most common and most pathogenic of these nematodes is *Ostertagia ostertagi*, which infects the abomasum of cattle. The disease syndrome caused by gastro-intestinal nematodes, commonly referred to as parasitic gastro-enteritis (PGE), drastically diminishes the economic viability of cattle production units (Kloosterman, A. et al., *Parasitology Today* 8, 330-335 (1992); 25 Vercruyse, J. and Claerebout, E., *Veterinary Parasitology* 98, 195-214 (2001)). The animals most at risk for PGE are calves during their first grazing season. Clinical PGE in grazing calves is characterized by (watery) diarrhea, weight loss, a dull hair coat, anorexia, a general loss of condition and eventually death (Anderson, N. et al., *Veterinary Record* 41, 196-204 (1965); Hilderson, H. et al., *Vlaams Diergeneeskundig Tijdschrift* 56, 30 269-29 (1987)). However, production losses are mainly due to sub-clinical infections, with no overt signs of disease. Substantial reductions in daily weight gain are observed in untreated first grazing season calves with sub-clinical infections (Shaw D.J., et al., *Veterinary Parasitology* 75, 115-131 (1998). Adult cows can still harbor large numbers of *O. ostertagi* (e.g. Borgsteede, F.H.M., et al., *Veterinary Parasitology* 89, 287-296 (2000); 35 Agneessens, J. et al., *Veterinary Parasitology* 90, 83-92 (2000)). Although gastrointestinal

nematode infections in adult cows are usually sub clinical, they are associated with decreased levels of milk production (Gross, S.J. et al., *Veterinary Record* 144, 581-587 (1999)). Carcass quality is also affected by gastrointestinal nematode infections, with reduced carcass weight, killing out percentage and related carcass measurements
5 (Entrocasso, C.M. et al., *Research in Veterinary Science* 40, 76-85 (1986)). Control of PGE in Europe is based almost exclusively on the use of anthelmintic drugs (Vercruyse, J. and Dorny, P., *International Journal for Parasitology* 29, 165-175 (1999)). However, the increased use of anthelmintics in cattle over the past two decades
10 (Borgsteede, F.H.M. et al., *Veterinary Parasitology* 78, 23-36 (1998); Schnieder, T. et al., *Veterinary Record* 145, 704-706 (1999); Claerebout, E. et al., *Vlaams Diergeneeskundig Tijdschrift* 69, 108-115 (2000)) has several drawbacks. The high costs of anthelmintic treatments, the negative effect of preventive anthelmintic treatments on the development of natural immunity against gastrointestinal nematodes (Vercruyse, J. et al., *Parasitology Today* 10, 129-132 (1994); Claerebout, E. and Vercruyse J., *Le Point Vétérinaire* 15 (*Numéro spécial*) 28, 175-179 (1997)), consumer concerns regarding drug residues in food products and in the environment (Wall, R. and Strong, L., *Nature* 327, 418-421 (1987); Steel, J.W. In: *NRA Special Review of Macroyclic Lactones*. National Registration Authority for Agricultural and Veterinary Chemicals, Canberra (1998); Strong, L., *Veterinary Parasitology* 48, 3-17 (1993)) and, last but not least, the increasing
20 incidence of parasite resistance against the available anthelmintics (Vermunt, J.J., et al., *Veterinary Record* 137, 43-45 (1995); Vermunt, J.J. et al., *New Zealand Veterinary Journal* 44, 188-193 (1996); Coles, G.C. et al., *Veterinary Record* 142, 255-256 (1998); Gill, J.H. and Lacey, E., *International Journal for Parasitology* 28, 863-877 (1998); and Fiel, C.A. et al., *Revista de Medicina Veterinaria* (Buenos Aires) 81, 310-315 (2000)) are
25 strong incentives for the producers to adopt alternative control systems (Vercruyse & Dorny (1999), *supra*). Vaccination is being considered as the most feasible solution (Knox, D.P., *Parasitology* 120, S43-S61 (2000)).
However, despite the evolution in biotechnology that allows the development of 'new generation' vaccines based on recombinant DNA technology, no vaccines against
30 gastrointestinal nematode parasites are available until now. The main problems that hamper the development of nematode vaccines in ruminants are (1) most parasite antigens that have been selected for vaccine development are 'hidden antigens', i.e. antigens that are not recognized by the host during a natural infection. Consequently, the immune response that is generated by vaccination with these antigens is not boosted by a
35 natural re-infection; (2) recombinant nematode proteins inducing a protective immune response have so far not been found.

It is an objective of the present invention to provide polypeptides that are capable of contributing to protection against the pathogenic effects of *Ostertagia ostertagi* infection in cattle.

5

It was now surprisingly found that 7 different polypeptides could be specifically identified and isolated, each of these different polypeptides being capable of inducing an immune response against *Ostertagia* parasites.

10 The inventors have found that these polypeptides can be used, either alone or in combination with each other, as vaccine components to provide a vaccine, which indeed contributes to the protection against *Ostertagia ostertagi* infection in cattle and helps to decrease the damage caused by *Ostertagia ostertagi*.

15 Three different approaches have been used for the detection of the genes encoding the vaccine components according to the invention. One approach, presented in detail under Example 1, uses specifically prepared anti-excretory-secretory protein rabbit antiserum for the detection of genes encoding immunoreactive *Ostertagia ostertagi* proteins. This approach has led to the finding of five novel immunogenic proteins for which the coding sequences are depicted in SEQ ID NO: 1, 3, 5, 7 and 9 as given below.

20

The gene encoding one such protein has now been cloned and sequenced and a nucleic acid sequence of the gene that comprises immunogenic determinants is depicted in SEQ ID NO: 7. The full-length gene encodes a protein of about 1600 amino acids (as partially depicted in SEQ ID NO: 8) with a molecular mass of >= 200 kD.

25

It is well known in the art, that many different nucleic acid sequences can encode one and the same protein. This phenomenon is commonly known as wobble in the second and especially the third base of each triplet encoding an amino acid. This phenomenon can result in a heterology for two nucleic acid sequences still encoding the same protein.

30 Therefore, in principle, two nucleic acid sequences having a sequence homology as low as 70 % can still encode one and the same protein.

Thus, one form of a first embodiment of the present invention relates to a nucleic acid sequence encoding an *Ostertagia ostertagi* protein or a part of said nucleic acid sequence 35 that encodes an immunogenic fragment of said protein wherein said nucleic acid

sequence or said part thereof has at least 85 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 7

The concept of immunogenic fragments is defined below. The length of a nucleic acid

- 5 sequence encoding an immunogenic fragment is usually at least 21 nucleotides, but preferably 24, 27, 30, 33 or even 36 nucleotides.

The molecular weight of all proteins according to the invention is determined in gel electrophoresis on a polyacrylamide gel. Due to slight variability of molecular weight

- 10 determination frequently encountered in the art, the molecular weight can vary. Therefore the molecular weight of the proteins according to the invention should be interpreted as to be its theoretical molecular weight +/- 5 kD.

Preferably, a nucleic acid sequence according to the invention encoding this *Ostertagia*

- 15 *ostertagi* protein or a part of that nucleic acid sequence that encodes an immunogenic fragment of that protein has at least 90 %, preferably 93 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 7

- 20 Even more preferred is a homology level of 98%, 99% or even 100%.

The level of nucleotide homology can be determined with the computer program "BLAST 2 SEQUENCES" by selecting sub-program: "BLASTN" that can be found at www.ncbi.nlm.nih.gov/blast/bl2seq/bl2.html.

- 25 A reference for this program is Tatiana A. Tatusova, Thomas L. Madden, *FEMS Microbiol. Letters* 174, 247-250 (1999). Parameters used are the default parameters:
Reward for a match: +1. Penalty for a mismatch: -2. Open gap: 5. Extension gap: 2. Gap x_dropoff: 50.

- 30 Nucleotide sequences that are complementary to the sequence depicted in SEQ ID NO: 1, 3, 5, 7, 9, 11 or 13, described herein, or nucleotide sequences that comprise tandem arrays of the sequences according to the invention, are also within the scope of the invention.

- 35 Another form of this embodiment relates to a nucleic acid sequence encoding a 28 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an

immunogenic fragment of said protein wherein said nucleic acid sequence or said part thereof has at least 85 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 3.

- 5 Preferably, a nucleic acid sequence according to the invention encoding this *Ostertagia ostertagi* protein or a part of that nucleic acid sequence that encodes an immunogenic fragment of that protein has at least 90 %, preferably 93 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 3.

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Even more preferred is a homology level of 98%, 99% or even 100%.

- Still another form of this embodiment relates to a nucleic acid sequence encoding a 25 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an 15 immunogenic fragment of said protein wherein said nucleic acid sequence or said part thereof has at least 85 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 5.

- Preferably, a nucleic acid sequence according to the invention encoding this *Ostertagia ostertagi* protein or a part of that nucleic acid sequence that encodes an immunogenic 20 fragment of that protein has at least 90 %, preferably 93 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 5.

- 25 Even more preferred is a homology level of 98%, 99% or even 100%.

- Again another form of this embodiment relates to a nucleic acid sequence encoding a 31 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein wherein said nucleic acid sequence or said part 30 thereof has at least 85 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 1.

- Preferably, a nucleic acid sequence according to the invention encoding this 31 kD *Ostertagia ostertagi* protein or a part of that nucleic acid sequence that encodes an 35 immunogenic fragment of that protein has at least 90 %, preferably 93 %, more preferably

95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 1.

Even more preferred is a homology level of 98%, 99% or even 100%.

5

Another form of this embodiment relates to a nucleic acid sequence encoding a 30 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein wherein said nucleic acid sequence or said part thereof has at least 85 % homology with the nucleic acid sequence of the 30 kD

10 *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 9.

Preferably, a nucleic acid sequence according to the invention encoding this 30 kD *Ostertagia ostertagi* protein or a part of that nucleic acid sequence that encodes an immunogenic fragment of that protein has at least 90 %, preferably 93 %, more preferably 15 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 9.

Even more preferred is a homology level of 98%, 99% or even 100%.

20 A second approach for the detection of vaccine components, presented in detail under Example 2, relied upon the analysis of components in a specific fraction of the parasite, the ES-fraction (excretory-secretory fraction) that play a role in establishing immunity against *Ostertagia ostertagi*. This approach surprisingly led to the finding of the 31 and 30 kD proteins described above (SEQ ID NO: 1 and 9). This provided a full confirmation of 25 the importance of the 31 and 30 kD proteins described above as vaccine components.

A third approach for the detection of vaccine components, presented in detail under Example 3, uses local antibodies obtained from mucus and Antibody Secreting Cell (ASC) culture supernatant. Although serum antibodies can in principle be used to screen for 30 candidate nematode antigens, local antibody responses produced at restricted tissue sites are not always detectable in serum. In addition, the persistence of serum antibodies makes it difficult to differentiate between previous and recent exposures to a pathogen. In contrast, local antibodies from the abomasal draining lymph nodes and from the mucus covering the abomasal mucosa are more specific for antigens present in the infected 35 tissue at the time of examination. It was shown in studies in rats and sheep that cell cultures, containing antibody secreting cells (ASC) induced *in vivo* in lymph nodes

draining the infected tissues, produce antibodies (ASC-probes) in the culture supernatant that specifically reflect the antigen exposure of the draining area and that stage-specific antigens are detected more readily by lymph node ASC-probes than by serum antibodies. Not only the draining lymph nodes but also the covering mucus-layer from the abomasum
5 are a source of local antibodies. After challenge infection of calves with *O. ostertagi*, a negative correlation between fecundity of the worm and parasite specific IgA in the mucus was observed (Claerebout, E. et al., 17th International Conference of the World Association for the Advancement of Veterinary Parasitology, Copenhagen, 1999). cDNA libraries of the 3 different parasitic stages were screened with the same antibody
10 probes to identify the nucleotide sequences that code for these antigens.

Details on the isolation of the genes encoding these antigens, and characterization of the protein antigens are presented in Examples 4 and 5.

This highly specific approach has been used for the selection of proteins and genes
15 encoding these proteins that can be directly linked to immune status instead of mere infected status. This approach has surprisingly revealed two more immunogenic proteins, for which the coding sequences are depicted below under SEQ ID NO: 11 and 13.

Therefore, another form of this embodiment relates to a nucleic acid sequence encoding a
20 24 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein wherein said nucleic acid sequence or said part thereof has at least 85 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 11.

25 Preferably, a nucleic acid sequence according to the invention encoding this *Ostertagia ostertagi* protein or a part of that nucleic acid sequence that encodes an immunogenic fragment of that protein has at least 90 %, preferably 93 %, more preferably 95 % homology with the nucleic acid sequence of the 24 kD *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 11.

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Even more preferred is a homology level of 98%, 99% or even 100%.

Again another form of this embodiment relates to a nucleic acid sequence encoding a 65 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein wherein said nucleic acid sequence or said part
35

thereof has at least 85 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 13.

Preferably, a nucleic acid sequence according to the invention encoding this *Ostertagia*

5 *ostertagi* protein or a part of that nucleic acid sequence that encodes an immunogenic fragment of that protein has at least 90 %, preferably 93 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 13.

10 Even more preferred is a homology level of 98%, 99% or even 100%.

Since the present invention discloses nucleic acid sequences encoding novel *Ostertagia ostertagi* proteins, it is now for the first time possible to obtain these proteins in sufficient quantities. This can e.g. be done by using expression systems to express the whole or 15 parts of the genes encoding the proteins or immunogenic fragments thereof according to the invention.

Therefore, in a more preferred form of this embodiment, the invention relates to DNA fragments comprising a nucleic acid sequence according to the invention. A DNA fragment is a stretch of nucleotides that functions as a carrier for a nucleic acid sequence 20 according to the invention. Such DNA fragments can e.g. be plasmids, into which a nucleic acid sequence according to the invention is cloned. Such DNA fragments are e.g. useful for enhancing the amount of DNA for use as a primer and for expression of a nucleic acid sequence according to the invention, as described below.

25 An essential requirement for the expression of the nucleic acid sequence is an adequate promoter functionally linked to the nucleic acid sequence, so that the nucleic acid sequence is under the control of the promoter. It is obvious to those skilled in the art that the choice of a promoter extends to any eukaryotic, prokaryotic or viral promoter capable of directing gene transcription in cells used as host cells for protein expression.

30 Therefore, an even more preferred form of this embodiment relates to a recombinant DNA molecule comprising a DNA fragment and/or a nucleic acid sequence according to the invention wherein the nucleic acid sequence according to the invention is placed under the control of a functionally linked promoter. This can be obtained by means of e.g. standard molecular biology techniques, e.g. Sambrook & Russell: "Molecular cloning: a 35 laboratory manual" (2001), Cold Spring Harbor Laboratory Press; ISBN: 0879695773.

Functionally linked promoters are promoters that are capable of controlling the transcription of the nucleic acid sequences to which they are linked.

Such a promoter can be the native promoter of a novel gene according to the invention or another promoter of *Ostertagia ostertagi*, provided that that promoter is functional in the

5 cells used for expression. It can also be a heterologous promoter. When the host cells are bacteria, useful expression control sequences, which may be used, include the Trp promoter and operator (Goeddel, et al., *Nucl. Acids Res.*, 8, 4057 (1980)); the lac promoter and operator (Chang, et al., *Nature*, 275, 615 (1978)); the outer membrane protein promoter (Nakamura, K. and Inouge, M., *EMBO J.*, 1, 771-775 (1982)); the
10 bacteriophage lambda promoters and operators (Remaut, E. et al., *Nucl. Acids Res.*, 11, 4677-4688 (1983)); the α -amylase (*B. subtilis*) promoter and operator, termination sequences and other expression enhancement and control sequences compatible with the selected host cell.

When the host cell is yeast, useful expression control sequences include, e.g., α -mating

15 factor. For insect cells the polyhedrin or p10 promoters of baculoviruses can be used (Smith, G.E. et al., *Mol. Cell. Biol.* 3, 2156-2165 (1983)). When the host cell is of vertebrate origin illustrative useful expression control sequences include the (human) cytomegalovirus immediate early promoter (Seed, B. et al., *Nature* 329, 840-842 (1987); Fynan, E.F. et al., *PNAS USA* 90, 11478-11482 (1993); Ulmer, J.B. et al., *Science* 259,
20 1745-1748 (1993)), Rous sarcoma virus LTR (RSV), Gorman, C.M. et al., *PNAS USA* 79, 6777-6781 (1982); Fynan et al., supra; Ulmer et al., supra), the MPSV LTR (Stacey et al., *J. Virology* 50, 725-732 (1984)), SV40 immediate early promoter (Sprague J. et al., *J. Virology* 45, 773 (1983)), the SV-40 promoter (Berman, P.W. et al., *Science* 222, 524-527 (1983)), the metallothionein promoter (Brinster, R.L. et al., *Nature* 296, 39-42 (1982)), the
25 heat shock promoter (Voellmy et al., *PNAS USA* 82, 4949-53 (1985)), the major late promoter of Ad2 and the β -actin promoter (Tang et al., *Nature* 356, 152-154 (1992)). The regulatory sequences may also include terminator and poly-adenylation sequences.
Amongst the sequences that can be used are the well known bovine growth hormone poly-adenylation sequence, the SV40 poly-adenylation sequence, the human
30 cytomegalovirus terminator and poly-adenylation sequences.

Bacterial, yeast, fungal, insect and vertebrate cell expression systems are very frequently used systems. Such systems are well known in the art and generally available, e.g. commercially through Clontech Laboratories Inc. (4030 Fabian Way, Palo Alto, California
35 94303-4607, USA). Next to these expression systems, parasite-based expression systems are attractive expression systems. Such systems are e.g. described in the

French Patent Application with Publication number 2 714 074, and in US NTIS Publication No US 08/043109 (Hoffman, S. and Rogers, W.: Public. Date 1 December 1993).

A very attractive expression system for heterologous nematode gene expression is a
5 nematodal expression system based upon the worm *Caenorhabditis elegans*. A heterologous expression system for this nematode has been described by Redmond, D.L. et al., in *Molecular and Biochemical Parasitology* 112, 125-131 (2001). See also Hashmi, S. et al., in *Trends in Parasitology* 17, 387-393 (2001).

The genes according to the present invention can be fused immediately downstream of a
10 *C. elegans* cystein protease promoter, cpr-5, which has been shown recently to direct expression to the gut of *C. elegans* (Redmond et al., 2001) and cloned into the pGEX-vector. The slow growing DR96 unc76(e911) *C. elegans* mutant strain can be transformed by micro-injection of plasmid DNA into the distal arm of the hermaphrodite gonad. The plasmid DNA can e.g. be prepared using the Qiagen method. *Ostertagia* genes according
15 to the invention can be co-injected with the repair plasmid p76-16B. The p76-16B plasmid rescues the unc76 phenotype and allows transformants to be identified through reversion back to the wild type phenotype. Transformed lines in which the second and subsequent generations show the wild type phenotype will be maintained. The presence of the injected construct in transgenic worms can easily be verified by PCR analysis of single
20 worms with primers developed specifically for the DNA of interest (Kwa et al., *Journal of Molecular Biology* 246, 500-510. (1995)). Transgenic worms, rescued by p76-16B, grow more quickly than the unc76(e911) mutants and allow rapid accumulation of transgenic worm material. Because of its rapid life-cycle, transformants can be grown *in vitro* in large quantities. Somatic extracts of transgenic worms can be prepared by grinding the
25 nematodes in a mortar under liquid nitrogen and resuspending them in 0.05M PBS containing 2% TritonX-100®. Fusion proteins will be purified by affinity chromatography using a Glutathione Sepharose column.

A still even more preferred form of this embodiment of the invention relates to Live
30 Recombinant Carriers (LRCs) comprising a nucleic acid sequence encoding an *Ostertagia ostertagi* protein or an immunogenic fragment thereof according to the invention, a DNA fragment according to the invention or a recombinant DNA molecule according to the invention. These LRCs are microorganisms or viruses in which additional genetic information, in this case a nucleic acid sequence encoding an *Ostertagia ostertagi* protein
35 or an immunogenic fragment thereof according to the invention has been cloned. Cattle infected with such LRCs will produce an immunological response not only against the

immunogens of the carrier, but also against the immunogenic parts of the protein(s) for which the genetic code is additionally cloned into the LRC, such as e.g. one or more of the novel *Ostertagia ostertagi* proteins gene according to the invention.

- 5 As an example of bacterial LRCs, attenuated *Salmonella* strains known in the art can very attractively be used.

Also, live recombinant carrier parasites have i.a. been described by Vermeulen, A. N. (*Int. J. Parasitol.* 28, 1121-1130 (1998)).

Furthermore, LRC viruses may be used as a way of transporting the nucleic acid

- 10 sequence into a target cell. Live recombinant carrier viruses are also called vector viruses. Viruses often used as vectors are Vaccinia viruses (Panicali et al; *PNAS USA* 79, 4927 (1982), Herpesviruses (E.P.A. 0473210A2), and Retroviruses (Valerio, D. et al.; in Baum, S.J., Dicke, K.A., Lotzova, E. and Pluznik, D.H. (Eds.), *Experimental Haematology today - 1988*. Springer Verlag, New York: pp. 92-99 (1989)).

15

The technique of *in vivo* homologous recombination, well known in the art, can be used to introduce a recombinant nucleic acid sequence into the genome of a bacterium, parasite or virus of choice, capable of inducing expression of the inserted nucleic acid sequence according to the invention in the host animal.

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Finally another form of this embodiment of the invention relates to a host cell comprising a nucleic acid sequence encoding a protein according to the invention, a DNA fragment comprising such a nucleic acid sequence or a recombinant DNA molecule comprising such a nucleic acid sequence under the control of a functionally linked promoter. This 25 form also relates to a host cell containing a live recombinant carrier comprising a nucleic acid molecule encoding an *Ostertagia ostertagi* protein or an immunogenic fragment thereof according to the invention.

A host cell may be a cell of bacterial origin, e.g. *Escherichia coli*, *Bacillus subtilis* and *Lactobacillus* species, in combination with bacteria-based plasmids as pBR322, or

- 30 bacterial expression vectors as the pEX-, pET-, pGEX-series, or with bacteriophages. The host cell may also be of eukaryotic origin, e.g. yeast-cells in combination with yeast-specific vector molecules, or higher eukaryotic cells like insect cells (Luckow et al.; *Bio-technology* 6, 47-55 (1988)) in combination with vectors or recombinant baculoviruses, plant cells in combination with e.g. Ti-plasmid based vectors or plant viral vectors (Barton, 35 K.A. et al.; *Cell* 32, 1033 (1983), mammalian cells like Hela cells, Chinese Hamster Ovary

cells (CHO) or Crandell-Rees Feline Kidney-cells, also with appropriate vectors or recombinant viruses.

Also, the host may be a nematode such as *C. elegans*, as explained above.

- 5 Another embodiment of the invention relates to the novel *Ostertagia ostertagi* proteins and to immunogenic fragments thereof according to the invention.

The concept of immunogenic fragments will be defined below.

- 10 One form of this embodiment relates to an *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order or preference, to the amino acid sequence as depicted in SEQ ID NO: 8.

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Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

20 The immunogenic fragments of the *Ostertagia ostertagi* protein as depicted in SEQ ID NO: 2, 4, 6, 8, 10, 12 and 14 according to the invention as described herein, preferably have a length of at least 7, more preferably 10, 15, 20, 30 or even 40 amino acids, in that order of preference.

25 A still even more preferred form of this embodiment relates to this *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

30 Another form of this embodiment relates to a 28 kD *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order or preference, to the amino acid sequence as depicted in SEQ ID NO: 4.

35 Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

A still even more preferred form of this embodiment relates to a 28 kD *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

5 Still another form of this embodiment relates to a 25 kD *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order or preference, to the amino acid sequence as depicted in SEQ ID NO: 6.

10 Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

15 A still even more preferred form of this embodiment relates to a 25 kD *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

20 Again another form of this embodiment relates to a 31 kD *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order or preference, to the amino acid sequence as depicted in SEQ ID NO: 2.

25 Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

A still even more preferred form of this embodiment relates to a 31 kD *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

30 One other form of this embodiment relates to a 30 kD *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order or preference, to the amino acid sequence as
35 depicted in SEQ ID NO: 10.

Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

A still even more preferred form of this embodiment relates to a 30 kD *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

Again an other form of this embodiment relates to a 24 kD *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order of preference, to the amino acid sequence as depicted in SEQ ID NO: 12.

Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

A still even more preferred form of this embodiment relates to a 24 kD *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

Again another form of this embodiment relates to a 65 kD *Ostertagia ostertagi* protein and to immunogenic fragments thereof, wherein the protein or immunogenic fragments have a sequence homology of at least 90%, preferably however 92%, more preferably 94%, 95% or even 96% homology, in that order or preference, to the amino acid sequence as depicted in SEQ ID NO: 14.

Even more preferred is a homology level of 97%, 98%, 99% or even 100% in that order of preference.

A still even more preferred form of this embodiment relates to a 65 kD *Ostertagia ostertagi* protein and immunogenic fragments of said protein, encoded by a nucleic acid sequence according to the present invention.

The level of protein homology can be determined with the computer program "BLAST 2 SEQUENCES" by selecting sub-program: "BLASTP", that can be found at www.ncbi.nlm.nih.gov/blast/bl2seq/bl2.html.

A reference for this program is Tatiana A. Tatusova, Thomas L. Madden, *FEMS Microbiol. Letters* 174, 247-250 (1999). Matrix used: "blosum62". Parameters used are the default parameters: Open gap: 11. Extension gap: 1. Gap x_dropoff: 50.

- 5 It will be understood that, for the particular proteins embraced herein, natural variations can exist between individual *Ostertagia ostertagi* strains. These variations may be demonstrated by (an) amino acid difference(s) in the overall sequence or by deletions, substitutions, insertions, inversions or additions of (an) amino acid(s) in said sequence. Amino acid substitutions which do not essentially alter biological and immunological activities, have been described, e.g. by Neurath et al in *The Proteins*, Academic Press New York (1979). Amino acid replacements between related amino acids or replacements which have occurred frequently in evolution are, inter alia, Ser/Ala, Ser/Gly, Asp/Gly, Asp/Asn, Ile/Val (see Dayhof, M.D., *Atlas of protein sequence and structure*, Nat. Biomed. Res. Found., Washington D.C. (1978), vol. 5, suppl. 3). Other amino acid substitutions 10 include Asp/Glu, Thr/Ser, Ala/Gly, Ala/Thr, Ser/Asn, Ala/Val, Thr/Phe, Ala/Pro, Lys/Arg, Leu/Ile, Leu/Val and Ala/Glu. Based on this information, Lipman and Pearson developed a method for rapid and sensitive protein comparison (*Science* 227, 1435-1441 (1985)) and determining the functional similarity between homologous proteins. Such amino acid 15 substitutions of the exemplary embodiments of this invention, as well as variations having deletions and/or insertions are within the scope of the invention as long as the resulting 20 proteins retain their immune reactivity.
- This explains why *Ostertagia ostertagi* proteins according to the invention, when isolated from different field isolates, may have homology levels of about 70%, while still representing the same protein with the same immunological characteristics.
- 25 Those variations in the amino acid sequence of a certain protein according to the invention that still provide a protein capable of inducing an immune response against infection with *Ostertagia ostertagi* or at least against the clinical manifestations of the infection are considered as "not essentially influencing the immunogenicity".
- 30 When a protein is used for e.g. vaccination purposes or for raising antibodies, it is however not necessary to use the whole protein. It is also possible to use a fragment of that protein that is capable, as such or coupled to a carrier such as e.g. KLH, of inducing an immune response against that protein, a so-called immunogenic fragment. An "immunogenic fragment" is understood to be a fragment of the full-length protein that still 35 has retained its capability to induce an immune response in a vertebrate host, e.g. comprises a B- or T-cell epitope. Shortly, an immunogenic fragment is a fragment that is

capable of inducing an antigenic response against an *Ostertagia ostertagi* protein according to the invention. At this moment, a variety of techniques are available to easily identify DNA fragments encoding antigenic fragments (determinants). The method described by Geysen et al. (Patent Application WO 84/03564, Patent Application WO 86/06487, US Patent nr. 4,833,092, *PNAS USA* 81, 3998-4002 (1984), *J. Imm. Meth.* 102, 259-274 (1987), the so-called PEPSCAN method is an easy to perform, quick and well-established method for the detection of epitopes; the immunologically important regions of the protein. The method is used worldwide and as such well known to man skilled in the art. This (empirical) method is especially suitable for the detection of B-cell epitopes. Also, given the sequence of the gene encoding any protein, computer algorithms are able to designate specific protein fragments as the immunologically important epitopes on the basis of their sequential and/or structural agreement with epitopes that are now known. The determination of these regions is based on a combination of the hydrophilicity criteria according to Hopp and Woods (*PNAS USA* 78, 38248-3828 (1981)), and the secondary structure aspects according to Chou and Fasman (*Advances in Enzymology* 47, 45-148 (1987) and US Patent 4,554,101). T-cell epitopes can likewise be predicted from the sequence by computer with the aid of Berzofsky's amphiphilicity criterion (*Science* 235, 1059-1062 (1987) and US Patent application NTIS US 07/005,885). A condensed overview is found in: Shan Lu, on common principles: *Tibtech* 9, 238-242 (1991); Good et al., on Malaria epitopes: *Science* 235, 1059-1062 (1987); Lu, for a review: *Vaccine* 10, 3-7 (1992); and Berzofsky, for HIV-epitopes: *The FASEB Journal* 5, 2412-2418 (1991). An immunogenic fragment usually has a minimal length of 6, more commonly 7-8 amino acids, preferably more than 8, such as 9, 10, 12, 15 or even 20 or more amino acids. The nucleic acid sequences encoding such a fragment therefore have a length of at least 18, more commonly 24 and preferably 27, 30, 36, 45 or even 60 nucleic acids.

Therefore, one form of still another embodiment of the invention relates to vaccines for combating *Ostertagia ostertagi* infection, that comprise at least one *Ostertagia ostertagi* protein or immunogenic fragments thereof, according to the invention as described above together with a pharmaceutically acceptable carrier.

- Still another embodiment of the present invention relates to the *Ostertagia ostertagi* proteins according to the invention or immunogenic fragments thereof for use in a vaccine.
- Again another embodiment of the present invention relates to the use of a nucleic acid sequence, a DNA fragment, a recombinant DNA molecule, a live recombinant carrier, a

host cell or a protein or an immunogenic fragment thereof according to the invention for the manufacturing of a vaccine, more specifically a vaccine for combating *Ostertagia ostertagi* infection.

- 5 One way of making a vaccine according to the invention is by growing the nematode, followed by biochemical purification of an *Ostertagia ostertagi* protein or immunogenic fragments thereof, from the nematode or the supernatant. This is however a very time-consuming way of making the vaccine.
- 10 It is therefore much more convenient to use the expression products of a gene encoding an *Ostertagia ostertagi* protein or immunogenic fragments thereof, according to the invention in vaccines. This is possible for the first time now because the nucleic acid sequences of genes encoding 7 novel *Ostertagia ostertagi* proteins suitable as vaccine components is provided in the present invention.
- 15 Vaccines based upon the expression products of these genes can easily be made by admixing the protein according to the invention or immunogenic fragments thereof according to the invention with a pharmaceutically acceptable carrier as described below.
- 20 Alternatively, a vaccine according to the invention can comprise live recombinant carriers as described above, capable of expressing the protein according to the invention or immunogenic fragments thereof. Such vaccines, e.g. based upon a *Salmonella* carrier or a viral carrier e.g. a Herpesvirus vector have the advantage over subunit vaccines that they better mimic the natural way of infection of *Ostertagia ostertagi*. Moreover, their self-
25 propagation is an advantage since only low amounts of the recombinant carrier are necessary for immunization.

Vaccines can also be based upon host cells as described above that comprise the protein or immunogenic fragments thereof according to the invention.

- 30 All vaccines described above contribute to active vaccination, i.e. they trigger the host's defense system.
Alternatively, antibodies can be raised in e.g. rabbits or can be obtained from antibody-producing cell lines as described below. Such antibodies can then be administered to the
35 cow. This method of vaccination, passive vaccination, is the vaccination of choice when an animal is already infected, and there is no time to allow the natural immune response

to be triggered. It is also the preferred method for vaccinating animals that are prone to sudden high infection pressure. The administered antibodies against the protein according to the invention or immunogenic fragments thereof can in these cases interfere with *Ostertagia ostertagi*. This approach has the advantage that it decreases or stops

5 *Ostertagia ostertagi* development.

Therefore, one other form of this embodiment of the invention relates to a vaccine for combating *Ostertagia ostertagi* infection that comprises antibodies against an *Ostertagia ostertagi* protein according to the invention or an immunogenic fragment of that protein, and a pharmaceutically acceptable carrier.

10

Still another embodiment of this invention relates to antibodies against an *Ostertagia ostertagi* protein according to the invention or an immunogenic fragment of that protein.

Methods for large-scale production of antibodies according to the invention are also known in the art. Such methods rely on the cloning of (fragments of) the genetic information encoding the protein according to the invention in a filamentous phage for phage display. Such techniques are described i.a. at the "Antibody Engineering Page" under "filamentous phage display" at
<http://aximt1.imt.uni-marburg.de/~rek/aepphage.html>, and in review papers by Cortese,
15 R. et al., (1994) in *Trends in Biotechn.* 12, 262-267., by Clackson, T. & Wells, J.A. (1994) in *Trends in Biotechn.* 12, 173-183, by Marks, J.D. et al., (1992) in *J. Biol. Chem.* 267, 16007-16010, by Winter, G. et al., (1994) in *Annu. Rev. Immunol.* 12, 433-455, and by Little, M. et al., (1994) *Biotechn. Adv.* 12, 539-555. The phages are subsequently used to screen camelid expression libraries expressing camelid heavy chain antibodies.
20 (Muyldermans, S. and Lauwerys, M., *Journ. Molec. Recogn.* 12, 131-140 (1999) and Ghahroudi, M.A. et al., *FEBS Letters* 414, 512-526 (1997)). Cells from the library that express the desired antibodies can be replicated and subsequently be used for large-scale expression of antibodies.

25 30 Still another embodiment relates to a method for the preparation of a vaccine according to the invention that comprises the admixing of antibodies according to the invention and a pharmaceutically acceptable carrier.

An alternative and efficient way of vaccination is direct vaccination with DNA encoding the relevant antigen. Direct vaccination with DNA encoding proteins has been successful for many different proteins. (As reviewed in e.g. Donnelly et al., *The Immunologist* 2, 20-26

(1993)). In the field of anti-parasite vaccines, protection against e.g. *Plasmodium yoelii* has been obtained with DNA-vaccination with the *Plasmodium yoelii* circumsporozoite gene (Vaccine 12, 1529-1533 (1994)). Protection against *Leishmania major* has been obtained with DNA-vaccination with the *Leishmania major* surface glycoprotein gp63 gene 5 (Vaccine 12, 1534-1536 (1994)).

This way of vaccination is also attractive for the vaccination of cattle against *Ostertagia ostertagi* infection. Therefore, still other forms of this embodiment of the invention relate to vaccines comprising nucleic acid sequences encoding a protein according to the invention 10 or immunogenic fragments thereof, vaccines comprising DNA fragments that comprise such nucleic acid sequences or vaccines comprising recombinant DNA molecules according to the invention, and a pharmaceutically acceptable carrier.

Examples of DNA plasmids that are suitable for use in a DNA vaccine according to the 15 invention are conventional cloning or expression plasmids for bacterial, eukaryotic and yeast host cells, many of said plasmids being commercially available. Well-known examples of such plasmids are pBR322 and pcDNA3 (Invitrogen). The DNA fragments or recombinant DNA molecules according to the invention should be able to induce protein expression of the nucleotide sequences. The DNA fragments or recombinant DNA 20 molecules may comprise one or more nucleotide sequences according to the invention. In addition, the DNA fragments or recombinant DNA molecules may comprise other nucleotide sequences such as immune-stimulating oligonucleotides having unmethylated CpG di-nucleotides, or nucleotide sequences that code for other antigenic proteins or adjuvating cytokines.

25

The nucleotide sequence according to the present invention or the DNA plasmid comprising a nucleotide sequence according to the present invention, preferably operably linked to a transcriptional regulatory sequence, to be used in the vaccine according to the invention can be naked or can be packaged in a delivery system. Suitable delivery 30 systems are lipid vesicles, ISCOMs®, dendromers, niosomes, microparticles, especially chitosan-based microparticles, polysaccharide matrices and the like, (see further below) all well-known in the art. Also very suitable as delivery system are attenuated live bacteria such as *Salmonella* species, and attenuated live viruses such as Herpesvirus vectors, as mentioned above.

35

Still other forms of this embodiment relate to vaccines comprising recombinant DNA molecules according to the invention.

DNA vaccines can e.g. easily be administered through intradermal application such as by using a needle-less injector. This way of administration delivers the DNA directly into the cells of the animal to be vaccinated. Amounts of DNA in the range between 10 pg and 1000 µg provide good results. Especially if the DNA is self-replicating, minor amounts will suffice. Preferably, amounts in the microgram range between 1 and 100 µg are used.

- 5 In a further embodiment, the vaccine according to the present invention additionally comprises one or more antigens derived from cattle pathogenic organisms and viruses, antibodies against those antigens or genetic information encoding such antigens and/or a pharmaceutical component such as an antibiotic.
- 10 Of course, such antigens, antibodies against such antigens, or genetic information can be of *Ostertagia ostertagi* origin, such as e.g. another *Ostertagia ostertagi* antigen. It can also be an antigen, antibodies or genetic information selected from another cow pathogenic organism or virus. Such organisms and viruses are preferably selected from the group of Bovine Herpesvirus, Bovine Viral Diarrhea virus, Parainfluenza type 3 virus, Bovine Paramyxovirus, Foot and Mouth Disease virus, *Pasteurella haemolytica*, Bovine Respiratory Syncytial Virus, *Theileria* sp., *Babesia* sp., *Trypanosoma* sp., *Anaplasma* sp., *Neospora caninum*, *Staphylococcus aureus*, *Streptococcus agalactiae*, *Mycoplasma*, *E. coli*, *Enterobacter*, *Klebsiella*, *Citrobacter*, *Cryptosporidium*, *Salmonella* and *Streptococcus dysgalactiae*.
- 15 20 25 Vaccines based upon one or more of the *Ostertagia ostertagi* proteins according to the invention are also very suitable as marker vaccines. A marker vaccine is a vaccine that allows to discriminate between vaccinated and field-infected cows e.g. on the basis of a characteristic antibody panel, different from the antibody panel induced by wild type infection. A different antibody panel is induced e.g. when an immunogenic protein present on a wild type *Ostertagia* is not present in a vaccine: the host will then not make antibodies against that protein after vaccination. Thus, a vaccine based upon any of the *Ostertagia ostertagi* proteins according to the invention would only induce antibodies against that specific protein, whereas a vaccine based upon a live wild-type, live attenuated or inactivated whole *Ostertagia ostertagi* would induce antibodies against all or most of the nematodal proteins.

A simple ELISA test, having wells comprising any other *Ostertagia* protein except for the *Ostertagia ostertagi* proteins according to the present invention and wells comprising only one or more purified *Ostertagia ostertagi* proteins according to the invention suffices to test serum from cows and to tell if the cows are either vaccinated with the protein vaccine
5 according to the invention or suffered from *Ostertagia ostertagi* field infection.

All vaccines according to the present invention comprise a pharmaceutically acceptable carrier. A pharmaceutically acceptable carrier can be e.g. sterile water or a sterile physiological salt solution. In a more complex form the carrier can e.g. be a buffer.
10

Methods for the preparation of a vaccine comprise the admixing of a protein or an immunogenic fragment thereof, according to the invention and/or antibodies against that protein or an immunogenic fragment thereof, and/or a nucleic acid sequence and/or a DNA fragment, a recombinant DNA molecule, a live recombinant carrier or host cell
15 according to the invention, and a pharmaceutically acceptable carrier.

Vaccines according to the present invention may in a preferred presentation also contain an immunostimulatory substance, a so-called adjuvant. Adjuvants in general comprise substances that boost the immune response of the host in a non-specific manner. A
20 number of different adjuvants are known in the art. Examples of adjuvants frequently used in cow vaccines are muramylpeptides, lipopolysaccharides, several glucans and glycans and Carbopol® (a homopolymer).

The vaccine may also comprise a so-called "vehicle". A vehicle is a compound to which the protein adheres, without being covalently bound to it. Such vehicles are i.a. bio-
25 microcapsules, micro-algicates, liposomes and macrosols, all known in the art.

Microparticles, more specifically those based upon chitosan, especially for use in oral vaccination are very suitable as vaccine vehicles.

A special form of such a vehicle, in which the antigen is partially embedded in the vehicle, is the so-called ISCOM® (EP 109.942, EP 180.564, EP 242.380)

30 In addition, the vaccine may comprise one or more suitable surface-active compounds or emulsifiers, e.g. Span® or Tween®.

35 Antigens will preferably be combined with adjuvants that are readily available and that are registered for use in domestic animals, e.g. aluminum hydroxide, a Th2-like modulating adjuvant.

Two alternative approaches for antigen delivery are especially suitable for application of the vaccines according to the present invention:

- a. systemic immunization with the inclusion of adjuvantia modulating immune responses towards the mucosa, such as vitamin D3 (Van der Stede, Y., et al., *Vaccine* 19, 1870-1878 (2001)) or QuilA®, and
- b. direct delivery to the respiratory mucosa by inhalation of naked DNA (plasmid) (Vanrompay, D., et al., *Immunology* 103, 106-112 (2001)).

Addition of CpG oligonucleotide sequences inside or outside the plasmid is also preferred for improving protection (Van der Stede, Y., et al., *Vet. Immunol. Immunopathol.*, 86, 31-41 (2002)).

Often, the vaccine is mixed with stabilizers, e.g. to protect degradation-prone proteins from being degraded, to enhance the shelf-life of the vaccine, or to improve freeze-drying efficiency. Useful stabilizers are i.a. SPGA (Bovarnik et al; *J. Bacteriology* 59, 509 (1950)), carbohydrates e.g. sorbitol, mannitol, trehalose, starch, sucrose, dextran or glucose, proteins such as albumin or casein or degradation products thereof, and buffers, such as alkali metal phosphates.

In addition, the vaccine may be suspended in a physiologically acceptable diluent. It goes without saying, that other ways of adjuvating, adding vehicle compounds or diluents, emulsifying or stabilizing a protein are also embodied in the present invention.

Vaccines according to the invention that are based upon the protein according to the invention or immunogenic fragments thereof can very suitably be administered in amounts ranging between 1 and 100 micrograms of protein per animal, although smaller doses can in principle be used. A dose exceeding 100 micrograms will, although immunologically very suitable, be less attractive for commercial reasons.

Vaccines based upon live attenuated recombinant carriers, such as the LRC-viruses, parasites and bacteria described above can be administered in much lower doses, because they multiply themselves during the infection. Therefore, very suitable amounts would range between 10^3 and 10^9 CFU/PFU for both bacteria and viruses.

Vaccines according to the invention can be administered e.g. intradermally, subcutaneously, intramuscularly, intraperitoneally, intravenously, or at mucosal surfaces such as orally or intranasally.

For efficient protection against disease, a quick and correct diagnosis of *Ostertagia ostertagi* infection is important.

Therefore it is another objective of this invention to provide diagnostic tools suitable for the
5 detection of *Ostertagia ostertagi* infection.

The nucleic acid sequences, the proteins and the antibodies according to the invention
are also suitable for use in diagnostics.

10 Therefore, another embodiment of the invention relates to nucleic acid sequences,
proteins and antibodies according to the invention for use in diagnostics.

The nucleic acid sequences or fragments thereof according to the invention can be used
to detect the presence of *Ostertagia ostertagi* in cows. A sample taken from the
15 abomasums of cows infected with *Ostertagia ostertagi* will comprise nucleic acid material
derived from said parasite, including nucleic acid sequences encoding for the protein
according to the invention. These nucleic acid sequences will hybridize with a nucleic acid
sequence according to the invention. Suitable methods for the detection of nucleic acid
sequences that are reactive with the nucleic acid sequences of the present invention
20 include hybridization techniques including but not limited to PCR techniques and NASBA®
techniques. Thus the nucleic acid sequences according to the invention can be used to
prepare probes and primers for use in PCR and or NASBA techniques.

A diagnostic test kit for the detection of *Ostertagia ostertagi* may e.g. comprise tools to
enable the reaction of *Ostertagia* nucleic acid isolated from the cows to be tested with
25 these tools. Such tools are e.g. specific probes or (PCR-) primers, also referred to as
primer fragments, based upon the nucleic acid sequences according to the invention. If
genetic material of *Ostertagia ostertagi* is present in the animal, this will e.g. specifically
bind to specific PCR-primers and, e.g. after cDNA synthesis, will subsequently become
amplified in PCR-reaction. The PCR-reaction product can then easily be detected in DNA
30 gel electrophoresis.

Standard PCR-textbooks give methods for determining the length of the primers for
selective PCR-reactions with *Ostertagia ostertagi* DNA. Primer fragments with a
nucleotide sequence of at least 12 nucleotides are frequently used, but primers of more
than 15, more preferably 18 nucleotides are somewhat more selective. Especially primers
35 with a length of at least 20, preferably at least 30 nucleotides are very generally

applicable. PCR-techniques are extensively described in C. Dieffenbach & G. Dveksler: *PCR primers: a laboratory manual*, CSHL Press, ISBN 879694473 (1995)).

Nucleic acid sequences according to the invention or primers of those nucleic acid sequences having a length of at least 12, preferably 15, more preferably 18, even more 5 preferably 20, 22, 25, 30, 35 or 40 nucleotides in that order of preference, wherein the nucleic acid sequences or parts thereof have at least 70 % homology with the nucleic acid sequence as depicted in SEQ ID NO: 1, 3, 5, 7, 9, 11 or 13 are therefore also part of the invention. Primers are understood to have a length of at least 12 nucleotides and a homology of at least 70%, more preferably 80%, 85%, 90%, 95%, 98%, 99% or even 10 100%, in that order of preference, with the nucleic acid sequence as depicted in SEQ ID NO: 1, 3, 5, 7, 9, 11 or 13. Such nucleic acid sequences can be used as primer fragments in PCR-reactions in order to enhance the amount of DNA that they encode or in hybridization reactions. This allows the quick amplification or detection on blots of specific 15 nucleotide sequences for use as a diagnostic tool for e.g. the detection of *Ostertagia ostertagi* as indicated above.

Another test on genetic material is based upon *Ostertagia* material obtained from e.g. a swab, followed by classical DNA purification followed by classical hybridization with 20 radioactively or color-labeled primer fragments. Colour-labelled and radioactively labeled fragments are generally called detection means. Both PCR-reactions and hybridization reactions are well-known in the art and are i.a. described in Sambrook & Russell, supra 15

Thus, one embodiment of the invention relates to a diagnostic test kit for the detection of 25 *Ostertagia ostertagi* nucleic acid sequences. Such a test comprises a nucleic acid sequence according to the invention or a primer fragment thereof.

A diagnostic test kit based upon the detection of antigenic material of the specific 30 *Ostertagia ostertagi* proteins according to the invention and therefore suitable for the detection of *Ostertagia ostertagi* infection may i.a. comprise a standard ELISA test. In one example of such a test the walls of the wells of an ELISA plate are coated with antibodies directed against any of the proteins according to the invention. After incubation with the material to be tested, labeled anti- *Ostertagia ostertagi* antibodies are added to the wells. A color reaction then reveals the presence of antigenic material from *Ostertagia ostertagi*. Therefore, still another embodiment of the present invention relates to diagnostic test kits 35 for the detection of antigenic material of *Ostertagia ostertagi*. Such test kits comprise

antibodies against a protein according to the invention or a fragment thereof according to the invention.

A diagnostic test kit based upon the detection in serum of antibodies against a protein of
5 *Ostertagia ostertagi* according to the invention and therefore suitable for the detection of *Ostertagia ostertagi* infection may i.a. comprise a standard ELISA test. In such a test the walls of the wells of an ELISA plate can e.g. be coated with an *Ostertagia ostertagi* protein according to the invention. After incubation with the material to be tested, labeled anti-bodies against that protein are added to the wells. A color reaction then reveals the
10 presence of antibodies against *Ostertagia ostertagi*.

Therefore, still another embodiment of the present invention relates to diagnostic test kits for the detection of antibodies against *Ostertagia ostertagi*. Such test kits comprise an *Ostertagia ostertagi* protein according to the invention or a fragment thereof according to the invention.

15 The design of the immunoassay may vary. For example, the immunoassay may be based upon competition or direct reaction. Furthermore, protocols may use solid supports or may use cellular material. The detection of the antibody-antigen complex may involve the use of labeled antibodies; the labels may be, for example, enzymes, fluorescent-,
20 chemoluminescent-, radio-active- or dye molecules.

Suitable methods for the detection of antibodies reactive with a protein according to the present invention in the sample include the enzyme-linked immunosorbent assay (ELISA), immunofluorescence test (IFT) and Western blot analyses.

25 The proteins or immunogenic fragments thereof according to the invention e.g. expressed as indicated above can be used to produce antibodies, which may be polyclonal, monospecific or monoclonal (or derivatives thereof). If polyclonal antibodies are desired, techniques for producing and processing polyclonal sera are well known in the art (e.g. Mayer and Walter, eds. *Immunochemical Methods in Cell and Molecular Biology*,
30 Academic Press, London (1987)).

Monoclonal antibodies, reactive against the protein according to the invention or an immunogenic fragment thereof according to the present invention, can be prepared by immunizing inbred mice by techniques also known in the art (Kohler and Milstein, *Nature*, 256, 495-497 (1975)).

EXAMPLES

Example 1

5 1.1. Parasite ES products, EX products and anti-ES rabbit serum preparation

EX products were prepared as described in Geldhof, P., et al., *Parasite Immunology* 24, 263-270 (2002). EX used in this example is comparable to S1 as described in this publication. Excretory-secretory products were prepared as described by Geldhof P, et al., *Parasitology* 121, 639-647 (2000). Rabbits were immunized three times, with one week 10 interval, with 100 µg of the obtained L₃, L₄ and Adult stage ES proteins in combination with Freund's adjuvant and bled three weeks after the last immunization. Polyclonal sera from these rabbits were used for immunoscreening of *O. ostertagi* cDNA libraries.

15 1.2. *O. ostertagi* cDNA library construction

Total RNA of L₃, L₄ and Adult parasites was prepared using TRIZOL® Reagent (GibcoBRL, Life Technologies). PolyA⁺ RNA was purified using mRNA Separator® Kit (Clontech Laboratories, Inc.). Three µg of mRNA was converted into first strand cDNA with random hexamer primers (SuperScript® Choice System for cDNA Synthesis, GibcoBRL, Life Technologies). Double stranded cDNA was modified with EcoRI-NotI 20 adapters and cloned into the lambda gt11 vector (Stratagene). Recombinant lambda phages were packaged (Gigapack®III Gold Packaging Extract, Stratagene) and the packaging reaction was titrated. The L₃ cDNA library was estimated to contain 1.15x10⁶ independent clones; the L₄ cDNA library 9.6x10⁶ and the Adult cDNA library contained 3.41x10⁶ plaque forming units. Upon amplification these cDNA libraries were 25 immunoscreened with the anti-ES rabbit sera.

30 1.3. Immunoscreening of cDNA library

Approximately 100,000 plaques were plated onto Luria Broth agar (8,000 plaques per plate) and replicas were made on nitrocellulose filters soaked in 10 mM isopropylthio-β-D-galactoside. Upon blocking the background (5% milk powder in PBST, Nestlé Gloria) the filters were incubated overnight with rabbit serum, diluted (1:200) in blocking buffer. Goat-anti-rabbit serum coupled to horseradish peroxidase (1:1000 dilution) was used as a conjugate and the antigen-antibody complexes were detected with diaminobenzidine. Reacting plaques were re-screened until a homogeneous population of immunopositive

recombinant phages was obtained. Purified plaques were resuspended in sterile SM buffer (50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 10 mM MgSO₄) and stored at 4°C.

1.4. Cloning and DNA sequence analysis of cDNA inserts

5 Phage inserts were PCR amplified with lambda gt11 primers:

λgt11F 5'- GGTGGCGACGACTCCTGGAGCCCG -3' (SEQ ID NO:15)

and

λgt11R 5'- TTGACACCAGACCAACTGGTAATG -3' (SEQ ID NO:16),

and cloned into a plasmid vector (pGEM-T®, Promega). DH5α *E. coli* transformants

10 containing the recombinant plasmid were selected on Luria Broth agar plates supplemented with 0.1 mg/ml ampicillin, 0.1mM isopropylthio-β-D-galactoside, and 40 µg/ml 5-bromo-4-chloro-3-indolyl-β-D-galactose and the cDNA inserts were PCR amplified with vector primers:

SP6 5'- ATTTAGGTGACACTATAGAA -3' (SEQ ID NO:17)

15 and

T7 5'- GTAATACGACTCACTATAGGGC -3' (SEQ ID NO:18).

The nucleotide sequence of the cDNA clones was determined by the dideoxy chain terminator method using fluorescent BigDye™ terminators in a 377 automated DNA sequencer (PE Biosystems). DNA sequence data were assembled (DNASTAR®, Inc.) 20 and compared with nucleic acid (Blast+Beauty) and amino acid sequences (BlastX+Beauty) in various databases (EMBL, GenBank, WU-Blast2 and Swiss-Prot).

Results of Example 1:

25

The screening method using specifically prepared anti-excretory-secretory rabbit antiserum for the detection of genes encoding immunoreactive *Ostertagia ostertagi* led to the detection of five novel genes encoding vaccine components.

All five genes were found to be present in the *Ostertagia ostertagi* Adult stage cDNA 30 library.

1) a gene encoding a novel immunogenic protein was found, of which the nucleotide sequence encoding important immunogenic determinants is given in SEQ ID NO: 7 The gene encodes a protein with a length of about 1600 amino acids and a molecular weight 35 of >= 200 kD. The amino acid sequence of an important immunoreactive part of this protein is given in SEQ ID NO: 8. As can be seen in Figure 1, several clones, one of which

is indicated by an arrow, comprise at least parts of the gene that encode an immunogenic part of this protein. It can be clearly seen that this protein is strongly recognized by antibodies against this protein.

- 5 2) a gene encoding a novel immunogenic protein of 28 kD was found. Most of the nucleotide sequence of this gene is given in SEQ ID NO: 3. The amino acid sequence of the protein is given in SEQ ID NO: 4. As can be seen in Figure 2B, in the lane denominated ES and EX (see under 1.1. for explanation) the clear band of about 28 kD representing this protein is strongly recognized by monospecific antisera purified on lanes 10 of plaque pure immunopositive clones encoding the protein.
- 15 3) a gene encoding a novel immunogenic protein of 25 kD was found. The nucleotide sequence of this gene is given in SEQ ID NO: 5. The amino acid sequence of the protein is given in SEQ ID NO: 6. As can be seen in Figure 2C, in the lane denominated EX (see 15 under 1.1. for explanation) the clear band of about 25 kD representing this protein is strongly and highly specifically recognized by monospecific antisera purified on lanes of plaque pure immunopositive clones encoding this protein.
- 20 4) a gene encoding a novel immunogenic protein of 31 kD was found. The nucleotide sequence of this gene is given in SEQ ID NO: 1. The amino acid sequence of the protein is given in SEQ ID NO: 2. In Figure 3B, in the boxed region, the four right-hand proteins are forms of this protein. (See also under results of Example 2). From Figure 3A it follows that the protein is strongly recognized by monospecific antisera purified on lanes of plaque 25 pure immunopositive clones encoding this protein.
- 30 5) a gene encoding a novel immunogenic protein of 30 kD was found. The nucleotide sequence of this gene is given in SEQ ID NO: 9. The amino acid sequence of the protein is given in SEQ ID NO: 10. In Figure 3B in the boxed region, the two left-hand proteins are forms of this protein. (See also under results of Example 2). From Figure 3A it follows that 30 the protein is strongly recognized by monospecific antisera purified on lanes of plaque pure immunopositive clones encoding this protein.

Example 2**2.1. Preparation of antigens**

Adult *O. ostertagi* parasites and Adult ES-products were obtained as described by

5 Geldhof et al. (2000, *Parasitology*, 121, 639-647).

2.2. Chromatography on Thiol-sepharose

Total ES was preincubated with a final concentration of 2.5 mM dithiothreitol (DTT) for 30 minutes at 37°C prior to chromatography. Excess DTT was removed by passage through

10 a 10 x 2.6 cm Sephadex® G-25 (Pharmacia) column and eluted with 10 mM Tris, 0.5 M NaCl, pH 7.4 at 5 ml/minute. An activated Thiol-Sepharose 4B (Sigma) column, 5 ml bed volume, was equilibrated in 10 mM Tris, 0.5 M NaCl, pH 7.4. Protein samples (10 mg/run) were applied to the Thiol-Sepharose 4B column at a flow rate of 5 ml/hour. Unbound material was eluted by washing the column with equilibration buffer (10 mM Tris, 0.5 M

15 NaCl, pH 7.4) till the OD₂₈₀ had returned to a steady baseline. Bound material was eluted with equilibration buffer containing 50 mM DTT at a flow rate of 5 ml/hour. The peak fractions were pooled. DTT was removed from the eluted proteins by passage, at 5 ml/minute, through a Sephadex® G-25 (Pharmacia) column in 10 mM Tris pH 7.4. The peak fractions were again pooled and protein content determined by the BCA method

20 (Pierce). Both purifications, S3- and ES-thiol, had a yield between 10 and 15 %. Aliquots of the ES-thiol fractions were removed for SDS-PAGE and substrate gel analysis. The remainder of the eluates was then stored at -70° C until required.

2.3. 1D and 2D gel electrophoresis

25 The peptide components of ES-thiol were visualized by Coomassie Blue staining (0.1 % Coomassie Blue R-250 in 40 % methanol and 10 % acetic acid) following fractionation of 10 µg protein sample by 10 % SDS-PAGE under reducing conditions.

The 2D gelectrophoresis was performed using the 1 PG-SDS/PAGE system according to Bjellqvist et al. (*Electrophoresis* 14, 1357-1365 (1993)). The protein samples were

30 precipitated by adding 10 volumes of ice-cold acetone and left for 2 hours at -20° C. The acetone was discarded after centrifugation. The pellet was resolved for 2 hours in rehydration solution containing 9 M urea, 4 % CHAPS (Pharmacia), Bromophenol Blue, 18 mM dithiothreitol and 2 % IPG buffer (Pharmacia). This sample, approximately 100 µg of protein, was loaded on 7 cm Immobiline strips (pH 3-10, Pharmacia) to perform the 35 isoelectric focusing. The strip was subsequently washed for 30 minutes in 50 mM Tris-Cl pH 8.8 containing 6 M Urea, 30 % glycerol (v/v), 2 % SDS (w/v), 64 mM dithiothreitol and

a trace of bromophenol blue. The second dimension was carried out on 12 % SDS-PAGE. Gels were stained by Coomassie Colloidal staining (Sigma).

2.4. Western blotting

5 The serum antibody responses of the calves to the immunizations with ES-thiol were evaluated by Western blotting using sera harvested one week after the second immunization. Five µg of ES-thiol was fractionated using 10 % SDS-PAGE under reducing conditions and then blot transferred onto a PVDF membrane. The blot sections were cut into strips and blocked overnight in 10 % horse serum in PBST. After 2 hours of probing
10 with pooled sera (diluted 1:400 in 2 % horse serum in PBST) from the different groups the conjugate (Rabbit anti-bovine-HPRO, Sigma, 1:8000 in 2 % horse serum in PBST) was added for one hour. Recognized antigens were visualized by adding 0.05 % 3,3-diaminobenzidine tetrachloride in PBS containing 0.01% H₂O₂ (v/v).

15 2.5. Mass spectrometric analysis

The mass spectrometric analysis was performed essentially as previously reviewed Jensen et al. (*Proteins*, Suppl 2, 74-89 (1998)). In short, protein spots were in-gel digested using trypsin and the peptides were subsequently purified with the AnchorChip® technology. The peptide samples were analyzed by MALDI-TOF mass spectrometry.
20 Remaining material was used for a LC-MS/MS analysis to determine the amino acid sequence of the different peptides.

Results of Example 2:

25

Peptide profile of ES-thiol and complete ES

Analysis of the ES-thiol protein fraction on 1D and 2D gel electrophoresis is shown in Figure 4. ES-thiol comprised a prominent band at ~30 kD as well as 3 lower molecular bands and around 6 peptides in the size range from 45 to 92 kD (Figure 3A). Analysis of
30 this protein fraction on 2D-gel is shown in Figure 3B. The prominent 30 kD band visible on the 1D gel migrates in approximately 6 spots between pI 5-7 on 2D-gel. Another 13 fainter spots with pI values ranging from 4 to 8 with molecular masses between 53 and 15 kD were visible in ES-thiol on 2D-gel (Figure 3B).

Antibody responses of immunized calves

The control animals showed some minor background recognition of a few peptides in ES-thiol (Figure 4). The ES-thiol group strongly recognized the 30 and the 31 kD antigen (Figure 4).

5

Mass-spectrometry results

The 6 abundant spots at 30 kD were excised from the gel and used in a MALDI-peptide mass fingerprint analysis (boxed in Figure 3B). Two different proteins were detected in these spots. The peptide mass fingerprint analysis indicated that spots number 3-6 10 contained the same 31 kD protein, as described above under 4) and spot 1 and 2 contained the 30 kD protein, as described above under 5). The remaining material was used in the LC-MS/MS analysis, which resulted in peptide sequences from spot 1-6. These showed 100% homology with previously characterized excretory-secretory antigen 15 as encoded by the genes encoding a 31 kD and 30 kD *Ostertagia ostertagi* protein, as described in Example 1, under 4) and 5).

Example 320 **3.1. Animals**

A total of 17 calves, male and female Holstein-cross breed, between 6 and 12 months old from 3 different farms received a natural infection with gastrointestinal nematodes during a first grazing season of at least 6 months.

To confirm the immune status of the calves, reductions in worm burdens were measured 25 after treatment at housing with benzimidazoles and subsequent challenge infection.

Calves of farm 1 (n=4) received a natural challenge during one month in the second grazing season (Claerebout et al., *Veterinary Parasitology* 75, 153-167 (1998)). Calves of farm 2 (n=6) and 3 (n=7) received an experimental challenge with 50,000 *O. ostertagi* L₃ larvae, one week after treatment. The *O. ostertagi* worm counts of these animals 30 ('immunized' animals) were compared with those from helminth free calves (n=6 for each farm), which received a similar challenge ('primary infected' animals). Reductions in worm counts were 48%, 45% and 24% for calves of farm 1, 2 and 3 respectively.

Sample collection3.2. Mucus collection

Abomasal mucus from all 17 'immunized' animals from the 3 different farms and from the

5 18 'primary infected' animals was collected by gently scraping the mucosal surface with a
glass microscope slide. Mucus scrapings were homogenized with an equal weight of
phosphate buffered saline (0.05 M PBS, pH 7.3, 3 mM Na-azide) using an Ultra-turrax
homogenizer (13,000 RPM, 3x1 min). The homogenates were centrifuged at 20,000g for
30 minutes. The supernatant was removed and stored at -70°C. To isolate the
10 immunoglobulins, the supernatant was treated with protein G-agarose beads (Roche).
Mucus (1 ml) was centrifuged (14,000 g, 4°C, 30 min) to remove the debris. 200 µl
Starting buffer (20 mM NaH₂PO₄, pH 7.0) were added to the supernatant to ensure that
the pH of the sample stayed neutral. After equilibration of the sample (2 washes with
starting buffer) 100 µl Protein G-agarose beads were added. The sample was placed on a
15 rotor for 2 h at 4°C to allow the binding of the Fc-parts of the Ig's to the beads.

Supernatant was collected and saved together with the first 5 washes (400 µl washing
buffer/wash, 20 mM NaH₂PO₄, 150 mM NaCl, 2 mM EDTA, pH 7.0). The bound Ig's were
eluted with 400 µl elution buffer (100 mM glycine, pH 2.7) until the OD of the elutions was
0. The fractions were immediately neutralized with 20% neutralization buffer (1 M Tris-
20 HCl, pH 9.0). The supernatant/wash fraction was again treated with protein G-agarose
beads to ensure that all antibodies present in the mucus sample were collected. The
treated mucus samples were pooled in 2 groups for each farm: the 'immunized' group and
the 'primary infected' group.

25 3.3. Antibody secreting cell probes (ASC-probes) collection

ACS-probes were collected from animals of farm 3 (n=13). Antibody secreting cell probes
(ASC-probes) designate the supernatant of a lymph node cell culture that was prepared
with the technique originally described by Meeusen and Brandon (*J. Immunol. Methods*
172, 71-76 (1994a); *Eur. J. Immunol.* 24, 469-474 (1994b)). In short, abomasal lymph
30 nodes were collected at necropsy and transported in cold PBS+1% penicillin-
streptomycin. Lymphocytes were harvested by cutting and teasing the nodes in 5 ml RPMI
medium (Gibco BRL), washed in RPMI medium and centrifuged (1,000 g, 10 min, 4°C).
The red blood cells were lysed by adding 20 ml lysis solution (2% Tris, pH 7.65, 0.8%
NH₄Cl), for 10 min with gentle shaking. Twenty ml RPMI containing 1% penicillin-
35 streptomycin and 2% horse serum was used to wash the cells 3 times. Cells were
resuspended to a final concentration of 5x10⁶ cells/ml in culture medium (RPMI

supplemented with 20 % horse serum, 1% penicillin-streptomycin, 1% sodium-pyruvate, 1% non-essential amino acids, 1% kanamycin, 0.1% gentamycin and 0.035% β -mercaptoethanol). Culture flasks containing 50 ml cell suspension were incubated at 37°C in an atmosphere of 5% CO₂ in air without stimulation. After 3 days, the cells were removed by 5 centrifugation (1000 g, 10 min) and 400 ml supernatant per animal were collected. The supernatant (ASC-probes) was concentrated 10 times in a SpeedVac® and pools of antibodies both from the 'immunized' animals and the 'primary infected' animals were made for screening Western Blots and cDNA libraries.

10 **3.4. cDNA library screening**

O. ostertagi L₃, L₄ and Adult cDNA libraries were constructed in λ gt11 phage, propagated on Y1090r⁻ cells and plated by standard methods (Sambrook & Russell, supra). Approximately 100,000 plaques of all 3 libraries were screened with ASC-probes and mucus antibodies. All plaques were first screened with a pool of antibodies of 'immunized' 15 animals from all three farms. All positive plaques were rescreened until a single plaque could be isolated. These positive plaques were rescreened with the antibody pool from 'primary infected' animals from all three farms. The plaques that were exclusively recognized by the antibodies from the 'immune' animals were retained, resuspended in 200 μ l of sterile SM buffer (50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 10 mM MgSO₄) and 20 stored at 4°C with a drop of chloroform. The others were designated false positives due to cross recognition of the antibodies from the 'primary infected' animals.

The inserts were amplified by PCR reaction with universal λ gt11 primers and the amplicon was gel-purified with a gel purification kit (Qiagen). The cDNA fragment was subcloned into pGEM-T vector (Promega) and transformed into DH5 α E. coli cells. Following blue-25 white screening (IPTG/X-gal) and PCR with SP6 and T7 vector primers, recombinant clones were selected and plasmid DNA was isolated using the Qiagen plasmid isolation kit. The nucleotide sequence of the cDNA clones was determined by the dideoxy chain terminator method using fluorescent BigDye™ terminators in a 377 automated DNA sequencer (PE Biosystems). Assembly and analysis of nucleotide and deduced amino 30 acid sequences were performed using the DNASTAR® software program.

Results of Example 3:

The screening method using local antibodies obtained from mucus and Antibody 35 Secreting Cell (ASC) culture supernatant made it possible that two additional novel genes encoding vaccine components were found:

1) a gene of 900 nucleotides was found in both the larval L₄ stage and in the Adult stage cDNA library. The nucleotide sequence of this gene is given in SEQ ID NO: 11. The gene encodes a protein with a length of 300 amino acids and a molecular weight of about 24 kD. The protein has an isoelectric point of pI 6.6. The amino acid sequence of the protein is given in SEQ ID NO: 12.

5 The arrows in Figure 5, left picture, show how bacteria expressing this protein are specifically recognized by antibodies found in the supernatant of lymph nodes isolated from immune animals. The importance of this finding is underlined by the fact that 10 antibodies isolated from primary infected animals do not at all react with these clones. This clearly indicates the importance of this protein in the induction of immunity. Further characterization of this protein is outlined in Example 4.

2) a gene of 1238 nucleotides was found in the larval L₃ stage and in the Adult stage 15 cDNA library. The nucleotide sequence of this gene is given in SEQ ID NO: 13. The gene encodes a protein of 65 kD. The amino acid sequence of the protein is given in SEQ ID NO: 14. The arrows in Figure 6, left picture, show how bacteria expressing this protein are specifically recognized by antibodies isolated from the mucus of immune animals. Again, the importance of this finding is underlined by the fact that antibodies isolated from 20 primary infected animals do not at all react with these clones. This clearly indicates the importance of this protein in the induction of immunity.

Details on the identification of the full-length gene are outlined in Example 5.

25 Example 4

4.1 Cloning of the gene for the 24 kD protein

A 653bp fragment was amplified from the gene clone encoding the 24 kD protein (De Maere et al., *Parasitology*, 125, 383-391 (2002)) by PCR using primers that also 30 incorporate restriction endonuclease sites (underlined). Primers used were:

24kForw 5'- GAATTCATGAAGTTGGTCGTG -3' (SEQ ID NO: 19)

and

24kRev 5'- CTCGAGTCAATAGATCCTTGTG -3' (SEQ ID NO: 20).

The PCR product was digested with restriction enzymes EcoRI and Xhol, gel-purified (Qiagen kit) and cloned in frame into the T7- /6xHisitidine-tagged vector pET21a (Novagen). The correct reading frame was confirmed by sequencing and the construct

was transformed into the BL21(DE3) strain of *Escherichia coli*. Recombinant protein expression was induced by addition of isopropyl- β thio galactosidase during 2h at 37°C.

Cells were centrifuged, resuspended in PBS and lysed by adding 0,1 volume of lysozyme. After a cycle of freezing (-70°C) and thawing, cell debris was spun down and 5 supernatant was collected. Cell debris was resuspended in the T7- BindBuffer® (+ 6M Ureum) for 1h on ice to resuspend the insoluble proteins.

Recombinant proteins were purified over a T7-tag affinity column and afterwards by a His-bind resin column.

10 **4.2 Polyclonal antibodies**

100 µg of recombinant protein was injected 3 times intramuscularly with 3 weeks interval in a rabbit. Pre-immune blood was taken just before the first immunization and the final bleeding was done 3 weeks after the last immunization.

15 **4.3 Sample collection**

Mucus collection and Antibody Secreting Cell Probe collection are described in example 3 above (sections 3.3 and 3.4), and in De Maere et al. (2002, supra).

4.4 Western Blotting

20 Recognition of native or recombinant 24 kD protein by ASC probes, Mucus antibodies or rabbit anti-24 kD protein serum was evaluated by Western blotting. Ten µg of *Ostertagia* extract or Excretion-Secretion product was fractionated using 10 % SDS-PAGE under reducing conditions and then transferred onto a PVDF membrane. The blot sections were cut into strips, blocked for 2 h in 10 % normal horse serum in PBST, and probed overnight 25 with ASC probes. Mucus antibodies or rabbit anti-24 kD protein serum and conjugate were then added: Rabbit anti-bovine-HPRO (H+L) (Jackson ImunoResearch Laboratories Inc.) at 1:8000, or HRPO- conjugated goat anti-rabbit (Sigma) at 1:6000 in 2 % normal horse serum in PBST. Strips were incubated for one hour. Recognized antigens were visualized by adding 0.05 % 3,3 diaminobenzidine tetrachloride in PBS containing 0.01% 30 H₂O₂ (v/v).

4.5 Quantitative RT PCR

RT-PCR was used to investigate transcripts of the gene encoding the 24 kD protein in *O. ostertagi* parasitic life stages. Three microgram of total RNA from each life stage (L₃, L₃-exsheathed, L₄ and Adult) was used for the cDNA synthesis using an oligo(dT) primer 35 (Superscript®, Life technologies). The oligonucleotide primers used for detection of transcripts

of the gene encoding the 24 kD protein were designed to amplify an approximately 300 basepair long cDNA. Actin (Oo-act), described by Vercauteren et al. (*Molecular and Biochemical Parasitology*, 126, 201-208 (2003)), was used as a constitutively expressed 'housekeeping' gene control to determine the uniformity of the reverse transcription reactions.

5 cDNA of the gene encoding the 24 kD protein was amplified and quantified using the Light Cycler® and the lightcycler-faststart DNA master SYBR green I kit (Roche, Mannheim, Germany). The reaction mixture consist of a master mix containing Taq DNA polymerase, dNTP mixture and SYBR green I, 2 mM MgCl₂, 5 pM of each primer and 2 µl of template cDNA in total of 20 µl. Confirmation of the specificity of the PCR-products was performed by
10 subjecting these products to a melting curve analysis, subsequent agarose gel electrophoresis and sequencing. The PCR analysis was performed in triplicate and quantification occurred using external standards of 24 kD protein and Oo-act cDNA. Calculation was performed with the Lightcycler analysis software. The relative amount of 24 kD protein expression was plotted as a ratio ((copy number of 24 kD protein / copy number of house keeping gene) x 10).

15

Results of Example 4:

Recombinant 24 kD protein (Figure 7A) was recognized by ASC-probes and Mucus
20 antibodies from immune animals (Figure 7B-C). This indicates that the epitopes of the recombinant protein resemble those of the native protein and that the recombinant protein has the same protective capacities as the native protein. Antibodies raised against the recombinant protein in rabbits, therefore also recognize the native protein on 1D gel (Figure 7D) and 2D gel (Figure 8A)

25 Antibodies to recombinant 24 kD protein were used to specify the stage specific expression of the protein on Western Blot (Figure 8B).

RT-PCR showed expression of the protein in all life stages, especially in the L₃ with sheath and in the L₄ stage (Figure 9A).

As the 24 kD protein is expressed predominantly in L₃ and L₄ stage larvae, a
30 vaccine based on this protein interferes with the development of the L₃ (the infective stage) and the L₄ larval parasite stages. Thereby reducing or preventing the establishment of an *Ostertagia* infection. This in turn leads to a reduced worm-load in the animal with all beneficial consequences set out herein.

Example 5**Obtaining the full length gene encoding the 65 kD protein**

Utilizing the Ad clone (De Maere et al., *Parasitology*, 125, 383-391 (2002)) as the basis for specific primer design, the complete sequence of the gene as depicted in SEQ ID NO: 13 was obtained by the technique of 5'/3'-Rapid Amplification of cDNA Ends (RACE). The 5'- RACE kit from GibcoBRL was employed to identify the 5' end of the gene for the 65 kD protein. First strand cDNA was produced in a reverse transcription reaction using the specific primer 65Rev1 (see below) on 2 µg Adult RNA. This cDNA was poly C tailed at its 3' end with terminal deoxytransferase and used as a template in a PCR with the Abridged Anchor Primer (AAP, GibcoBRL):

AAP: 5'- GGCCACGCGTCGACTAGTACGGGIIGGGIIGGGIIG -3' (SEQ ID NO: 21)
and gene specific primer 65Rev2 (see below). The 5' RACE PCR product was cloned and sequenced.

The 3'-RACE kit from GibcoBRL was employed to identify the 3' end of the gene for the 65 kD protein. Briefly, first strand cDNA was produced in a reverse transcription reaction using an oligo(dT)-containing Adapter Primer (AP) on 2 µg Adult RNA. This cDNA was used as a template in a PCR with a gene specific primer 65kForw (see below) and the Universal Amplification Primer (UAP, GibcoBRL):

UAP 5'- CUACUACUACUAGGCCACGCGTCGACTAGTAC -3' (SEQ ID NO: 22)

The 3' RACE PCR product was cloned and sequenced.

Alignment of all the sequence data made it possible to design new gene specific primers comprising the start- and stopcodon, For65 and Rev65 (see below). The SUPERSCRIPT™ Preamplification System for First Strand cDNA Synthesis (GibcoBRL) was used to create template for a PCR with these primers to obtain the full length cDNA.

25

Gene specific primers employed to identify the full length coding sequence of the cDNA for the 65 kD protein are:

65Rev1: 5'- CAGCAATGGATACCGAATGAC -3' (SEQ ID NO: 23)

65Rev2: 5'- AGTGACTTCATCATTGCTGGTG -3' (SEQ ID NO: 24)

30 65kForw: 5'- TGATGATGAAGAACGAGAGGA -3' (SEQ ID NO: 25)

For65: 5'- GGATCCATGAGGCTGATATTGCTCATTAA -3' (SEQ ID NO: 26)

Rev65: 5'- CTCGAGGCAGAGTCCACACGACTTTGG -3' (SEQ ID NO: 27)

Quantitative RT-PCR

RT-PCR was used to investigate transcripts of the gene for the 65 kD protein in the parasitic *O. ostertagi* life stages. Three microgram of total RNA from each life stage (L_3 , L_3 -exsheathed, L_4 and Adult) were used for the cDNA synthesis using an oligo(dT) primer (Superscript, Life 5 technologies). The oligonucleotide primers used for detection of the transcript for the 65 kD protein were designed to amplify an approximately 300-500 basepair long cDNA. Actin (Oo-act), described by Vercauteren et al. (2003, supra), was used as a constitutively expressed 'housekeeping' gene control to determine the uniformity of the reverse transcription reactions.

cDNA for the 65 kD protein was amplified and quantified as described in Example 4, 10 section 4.5.

Results of Example 5:

15 The complete coding sequence of the gene encoding the 65 kD protein, as depicted SEQ ID NO:13 is 1722 bp long and codes for a protein with a molecular weight of 65 kD (SEQ ID NO:14). The N-terminal aa sequence contains a putative signal-sequence that is probably cleaved between aa 16 and 17 (Glycine-Glycine). The encoded protein sequence contains 5 N-glycosylation sites, a zinc-binding region (cd00203: 20 HEXXHALGFXHEXXRXDR) and a pfam01400 domain (HEXXHXXG) this places it in the family of Astacin's (Peptidase family M12A).

RT-PCR revealed stage specific expression of the gene for the 65kD protein by *Ostertagia ostertagi*. Transcripts were detected in the L_3 and the Adult stages with particular higher expression level in the Adult life stage (Figure 9B). This is in conformity 25 with the screening of the cDNA library (De Maere et al., 2002, supra).

As the 65 kD protein is expressed predominantly in Adult stages, a vaccine based on this protein interferes with Adult parasite development. This leads to reduced production of eggs, which in turn reduces the contamination of the fields. This reduces the worm-burden and contamination levels later in the season.

Example 6**Expression in the baculovirus expressionvector system**

The coding regions for the 65, 28, 31, and 24 kD proteins of the invention were subcloned
5 from their respective vectors into a pFastBac® plasmid (Invitrogen) using standard techniques. These FastBac constructs were transfected into Sf9 insect cells, to produce recombinant baculoviruses, according to the manufacturer's instructions (Invitrogen). Next expression cultures were run, using Sf9 and Sf158 insect cells, which were cultured in microcarrier spinner flasks of 100 and 250 ml. Serum free culture media used were
10 CCM3™ (Hyclone), and SF900-II™ (Invitrogen). Cells were infected at an m.o.i. of 0.1-0.5 and cultured for 3-4 days. Then cultures were centrifuged, culture supernatant was harvested, and cell pellets were resuspended 10 x concentrated in PBS. Triton X-100® was added to all samples to a concentration of 0.2 % v/v. Samples were extracted
15 overnight at room temperature, centrifuged, and supernatants were stored at -20°C until use.

Extract-supernatants were run on standard SDS/PAGE gels alongside appropriate markers, blotted onto Immobilon-P® transfer membrane (Millipore), membranes were stained with anti His-tag monoclonal antibody (Sigma), and visualized.

Figure 10 shows the results of these baculovirus expressions in Western blot; the
20 proteins of the invention were most abundant in the insect cell-pellet samples. These have been used to formulate vaccines for vaccinations.

LEGEND TO THE FIGURES

Figure 1: Dot-blot of lysed bacteria comprising a nucleotide sequence encoding (at least an immunogenic part of) the protein as depicted in SEQ ID NO: 8. Screening was done
5 with specifically prepared anti-excretory-secretory rabbit antiserum (See Example 1). An arrow indicates one of the positive clones.

Figure 2: Western-blots; in panel 2B of the 28 kD protein with anti-ES and anti-EX rabbit
10 antiserum (see Example 1), and in panel 2C of the 25 kD protein with anti-ES and anti-EX rabbit antiserum (see Example 1).

Figure 3: Analysis of the ES-thiol protein fraction (see also Example 2); in panel 3A a 1D gel electrophoresis, and in panel 3B a 2D gel electrophoresis. The 2D gel shows the 31 kD protein (the four right-most spots in the boxed area) and the 30 kD protein (the two left-
15 most spots in the boxed area).

Figure 4: Antibody response of ES-thiol-immunized calves against ES-fraction proteins.

Figure 5: Dot-blot of lysed bacteria comprising a nucleotide sequence encoding the 24 kD
20 protein as depicted in SED ID NO: 12. Screening was done with specifically prepared antibodies from lymph node supernatant of immune animals (left-hand picture). (See also Examples 3, and 4). Arrows indicate some of the positive clones. The right-hand picture shows a comparable dot-blot, now incubated with antibodies of primary infected animals.
With these antibodies no positive clones are recognized.

25

Figure 6: Dot-blot of lysed bacteria comprising a nucleotide sequence encoding the 65 kD protein as depicted in SED ID NO: 14. Screening was done with specifically prepared antibodies from mucus of immune animals (left-hand picture). (See also Examples 3, and 5). Arrows indicate some of the positive clones. The right-hand picture shows a comparable dot-blot, now incubated with antibodies of primary infected animals. With these antibodies no positive clones are recognized.

30
35 Figure 7: Electrophoretic characterization of the 24 kD protein; in panel 7A: result of expression of recombinant 24 kD protein in *E. coli*; panel 7B: a Western blot of rec 24 kD protein developed with ASC probe antibodies from immune animals; in panel 7C: Western

blot of rec 24 kD protein developed with Mucus antibodies from immune animals, and panel 7D: Western Blot of L₄ extract developed with rabbit anti-24 kD protein antibodies.

Figure 8: Characterization of the 24 kD protein; in panel 8A: 2D gel electrophoresis and

5 Western Blotting of Adult extract from *Ostertagia ostertagi*, developed with specific antibodies against *E. coli* expressed recombinant 24 kD protein, raised in rabbits, and panel 8B: stage specific expression of 24 kD protein, developed with anti-rec 24 kD protein antibodies.

10 Figure 9: Results of quantitative RT-PCRs to detect stage specific expression; in panel 9A for 24 kD protein (see Example 4), and in panel 9B for the 65 kD protein (see Example 5).

Figure 10: Western blot of *Ostertagia* proteins expressed in the baculovirus expressionvector system (see Example 6), staining was with anti-His antibody

15 Lanes 1 and 11: BioRad Protein Precision Marker®

Lane 2: 65 kD protein, supernatant + 0.2% TX-100

Lane 3: 65 kD protein, cell-pellet in PBS + 0.2% TX-100 (5x conc.)

Lane 4: 28 kD protein, supernatant + 0.2% TX-100

Lane 5: 28 kD protein, cell-pellet in PBS + 0.2% TX-100 (5x conc.)

20 Lane 6: 31 kD protein, supernatant + 0.2% TX-100

Lane 7: 31 kD protein, cell-pellet in PBS + 0.2% TX-100 (5x conc.)

Lane 8: 24 kD protein, supernatant + 0.2% TX-100

Lane 9: 24 kD protein, cell-pellet in PBS + 0.2 % TX-100 (5x conc.)

Lane 10: empty

CLAIMS

1. Nucleic acid sequence encoding an *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 7.
2. Nucleic acid sequence encoding a 28 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 3.
3. Nucleic acid sequence encoding a 25 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 5.
4. Nucleic acid sequence encoding a 31 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 1.
5. Nucleic acid sequence encoding a 30 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 9.

6. Nucleic acid sequence encoding a 24 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 11.
7. Nucleic acid sequence encoding a 65 kD *Ostertagia ostertagi* protein or a part of said nucleic acid sequence that encodes an immunogenic fragment of said protein, said nucleic acid sequence or said part thereof having at least 85%, preferably 90 %, more preferably 95 % homology with the nucleic acid sequence of the *Ostertagia ostertagi* protein gene as depicted in SEQ ID NO: 13.
8. DNA fragment comprising a nucleic acid sequence according to claim 1-7.
9. Recombinant DNA molecule comprising a nucleic acid sequence according to claim 1-7 or a DNA fragment according to claim 8, under the control of a functionally linked promoter.
10. Live recombinant carrier comprising a nucleic acid sequence according to claim 1-7, a DNA fragment according to claim 8 or a recombinant DNA molecule according to claim 9.
11. Host cell comprising a nucleic acid sequence according to claim 1-7, a DNA fragment according to claim 8, a recombinant DNA molecule according to claim 9 or a live recombinant carrier according to claim 10.
12. An *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 8.
13. A 28 kD *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 4.

14. A 25 kD *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 6.
15. A 31 kD *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 2.
16. A 30 kD *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 10.
17. A 24 kD *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 12.
18. A 65 kD *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, characterized in that said protein or immunogenic fragment thereof has a sequence homology of at least 90%, preferably 92 %, more preferably 94% to the amino acid sequence as depicted in SEQ ID NO: 14.
19. An *Ostertagia ostertagi* protein or an immunogenic fragment of said protein, according to claim 12-18, characterized in that said protein or immunogenic fragment is encoded by a nucleic acid sequence according to claim 1-7.
20. An *Ostertagia ostertagi* protein or an immunogenic fragment thereof, according to claim 12-19 for use in a vaccine.

21. Use of a nucleic acid sequence according to claim 1-7, a DNA fragment according to claim 8, a recombinant DNA molecule according to claim 9, a live recombinant carrier according to claim 10, a host cell according to claim 11 or a protein according to claim 12-19 or an immunogenic fragment thereof for the manufacturing of a vaccine for combating *Ostertagia ostertagi* infection.
22. Vaccine for combating *Ostertagia ostertagi* infection, characterized in that said vaccine comprises at least one *Ostertagia ostertagi* protein or an immunogenic fragment of said protein according to claim 12-19 and a pharmaceutically acceptable carrier.
23. Vaccine for combating *Ostertagia ostertagi* infection, characterized in that said vaccine comprises a nucleic acid sequence according to claim 1-7, a DNA fragment according to claim 8, a recombinant DNA molecule according to claim 9, a live recombinant carrier according to claim 10 or a host cell according to claim 11 and a pharmaceutically acceptable carrier.
24. Vaccine for combating *Ostertagia ostertagi* infection, characterized in that said vaccine comprises antibodies against a protein or an immunogenic fragment thereof according to claim 12-19 and a pharmaceutically acceptable carrier.
25. Vaccine according to claim 22-24, characterized in that said vaccine comprises an adjuvant.
26. Vaccine according to claim 22-25, characterized in that said vaccine comprises an additional antigen derived from a virus or micro-organism pathogenic to cattle, an antibody against said antigen or genetic information encoding said antigen and/or a pharmaceutical component.
27. Vaccine according to claim 26, characterized in that said virus or micro-organism pathogenic to cattle is selected from the group of Bovine Herpesvirus, Bovine Viral Diarrhea virus, Parainfluenza type 3 virus, Bovine Paramyxovirus, Foot and Mouth Disease virus, *Pasteurella haemolytica*, Bovine Respiratory Syncytial Virus, *Theileria* sp., *Babesia* sp., *Trypanosoma* species, *Anaplasma* sp., *Neospora caninum*, *Staphylococcus aureus*, *Streptococcus agalactiae*, *Mycoplasma*, *E. coli*, *Enterobacter*, *Klebsiella*, *Citrobacter*, *Cryptosporidium*, *Salmonella* and *Streptococcus dysgalactiae*.

28. Method for the preparation of a vaccine according to claim 22-27, said method comprising the admixing of a nucleic acid sequence according to claim 1-7, a DNA fragment according to claim 8, a recombinant DNA molecule according to claim 9, a live recombinant carrier according to claim 10, a host cell according to claim 11, a protein according to claim 12-19 or antibodies against a protein according to claim 12-19, and a pharmaceutically acceptable carrier.
29. A diagnostic kit comprising suitable detection means and a nucleic acid sequence according to claim 1-7 or a primer fragment thereof, or a protein according to claim 12-19, or an immunogenic fragment of said protein, or antibodies that are reactive with a protein according to claim 12-19.

FIGURES

Figure 1:

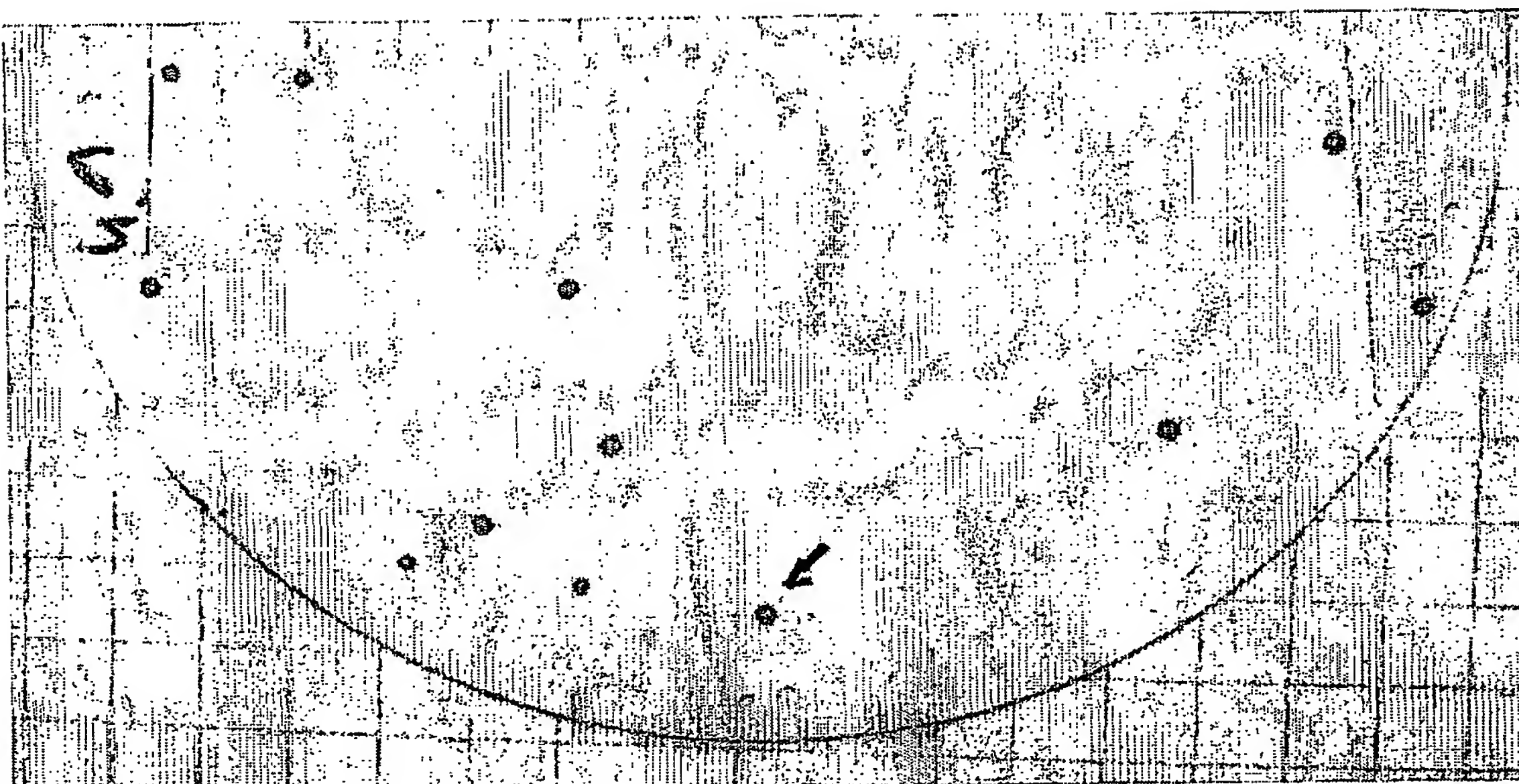


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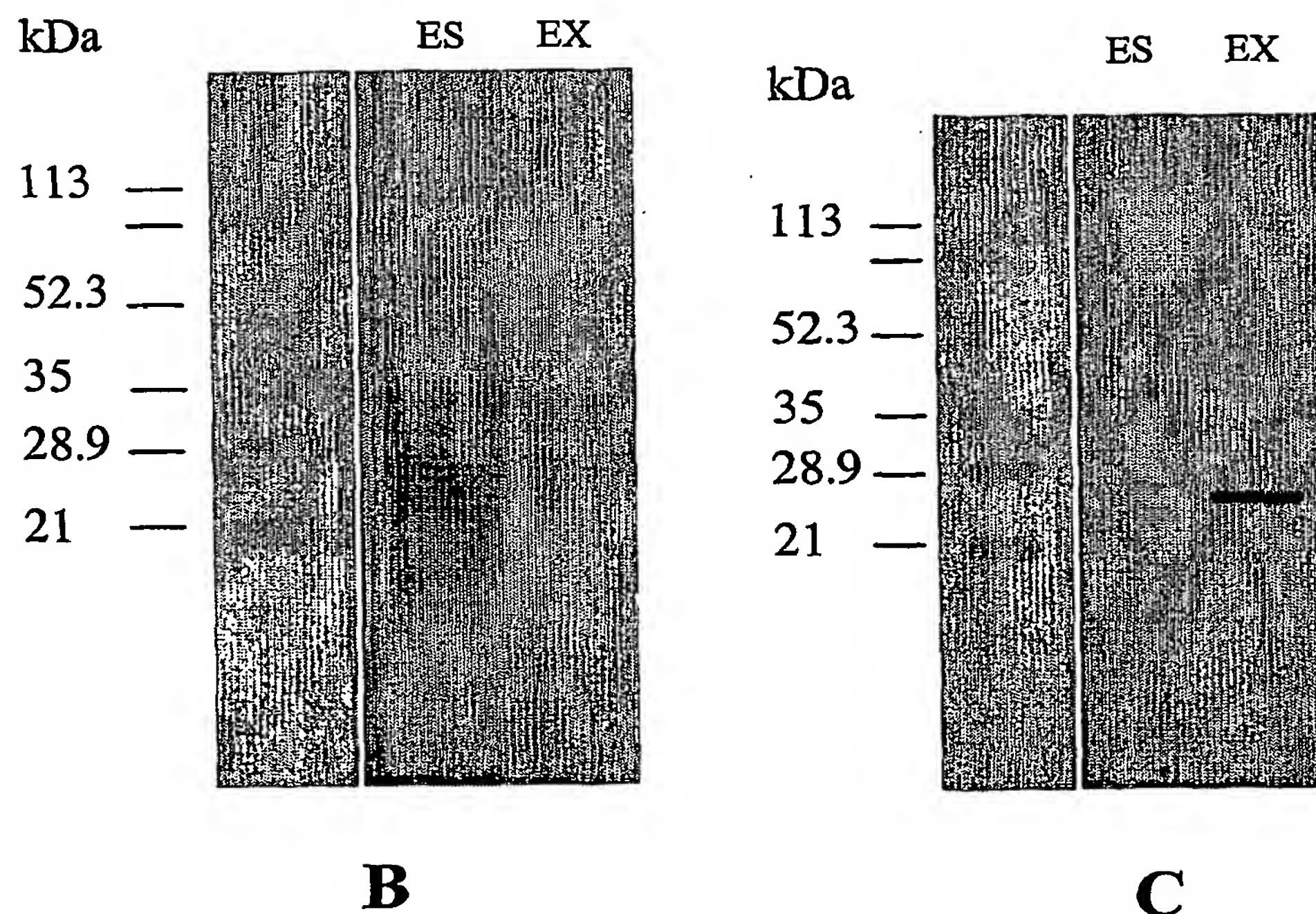


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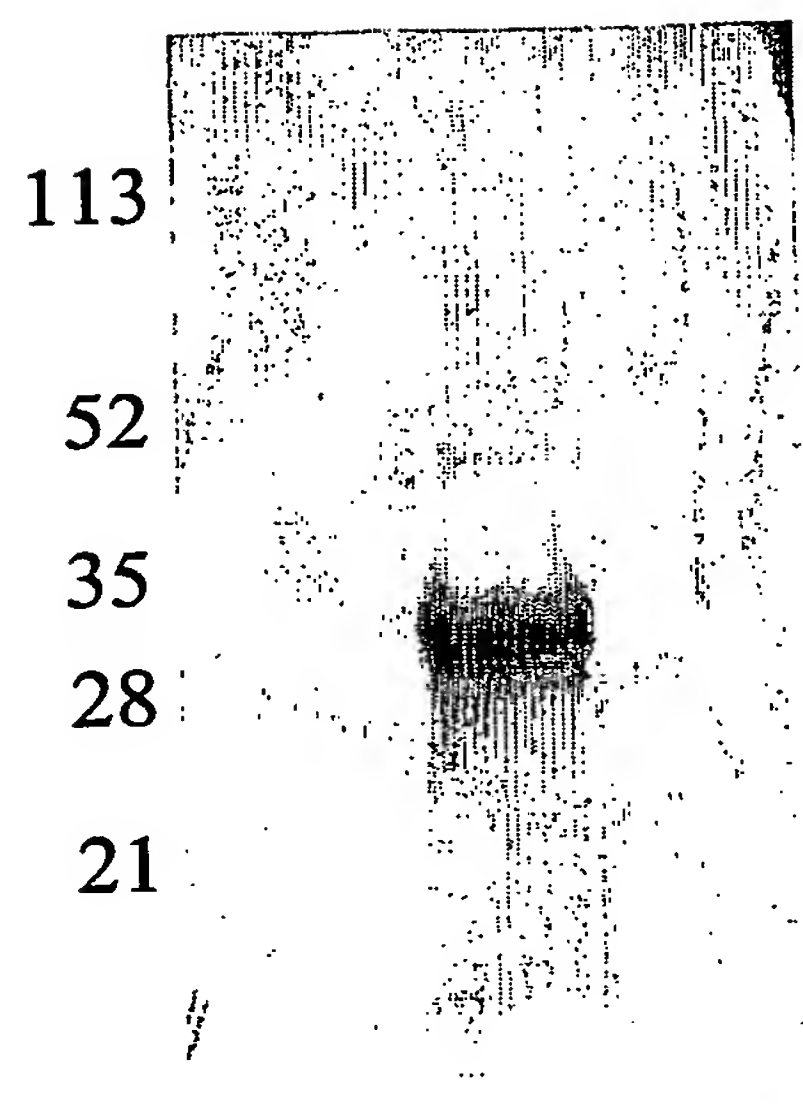


Figure 3 B

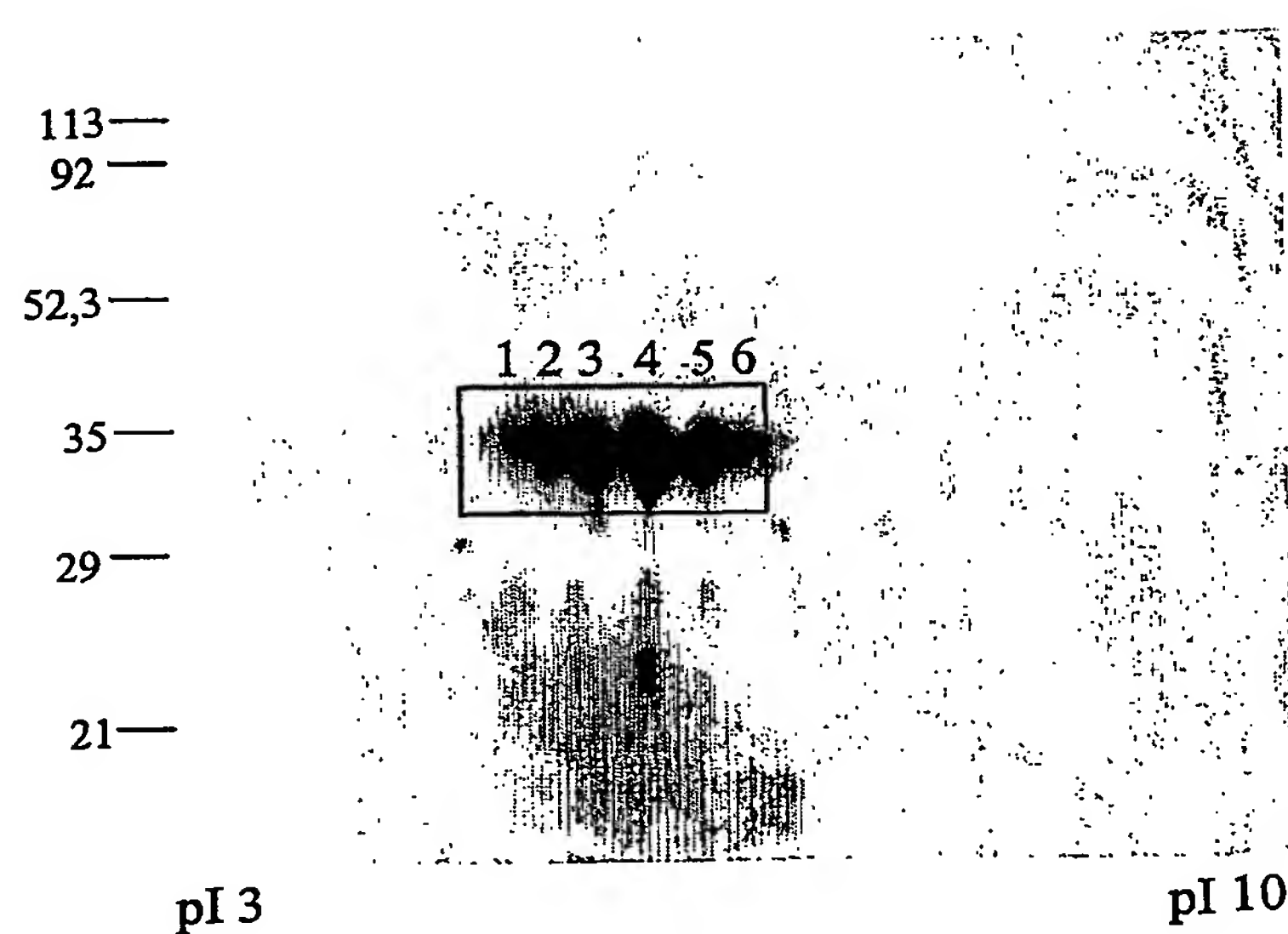


Figure 4

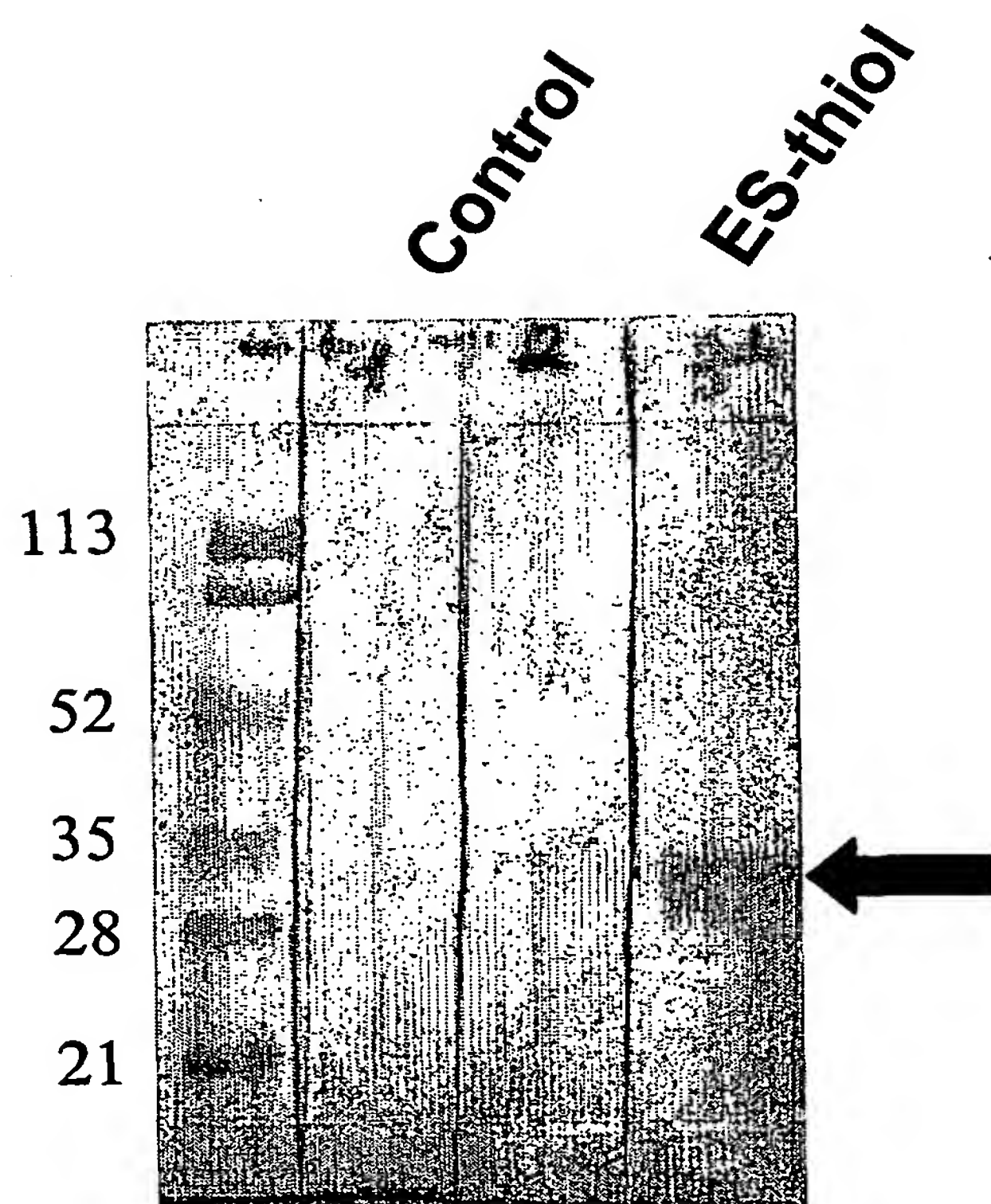


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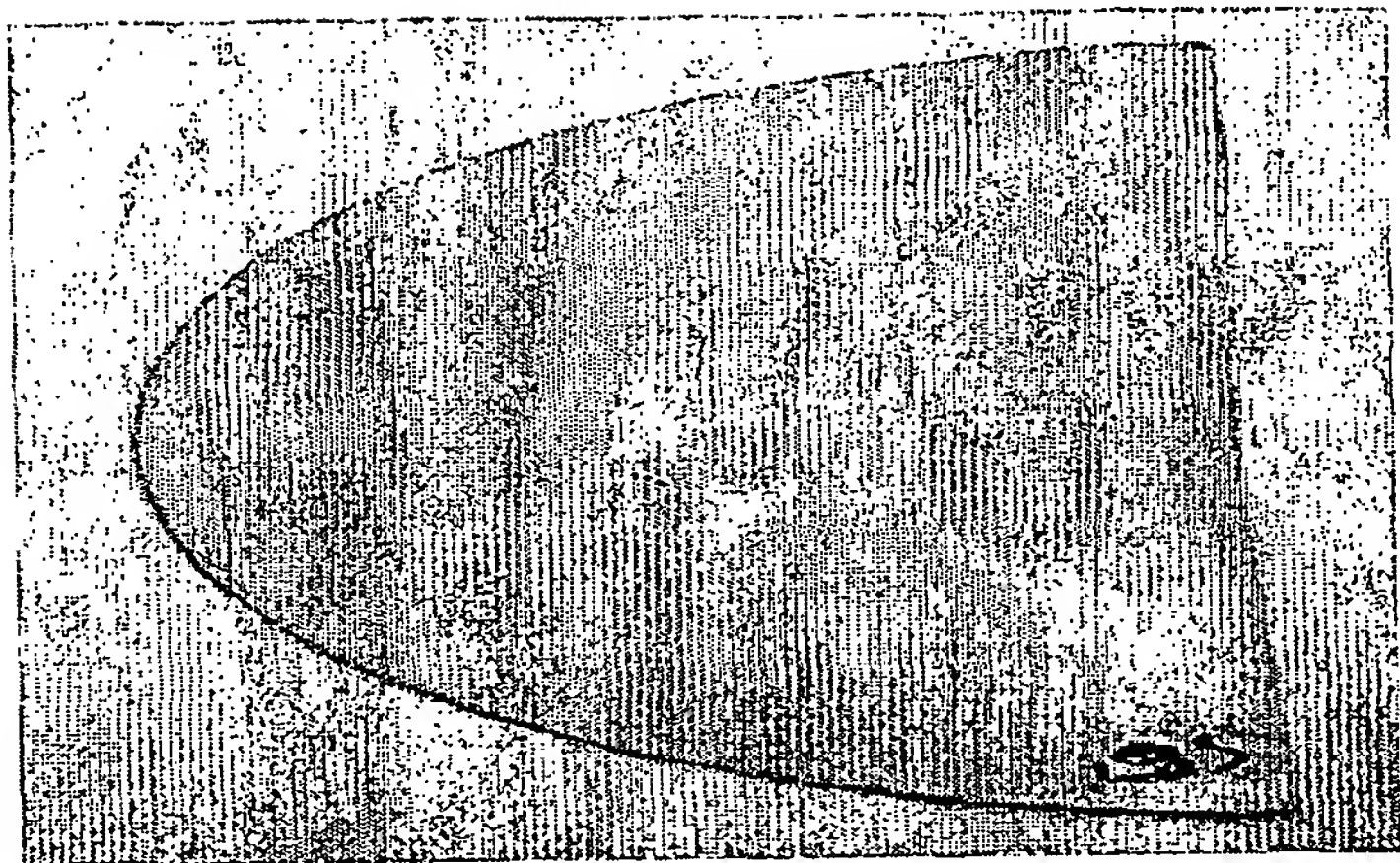


Figure 6

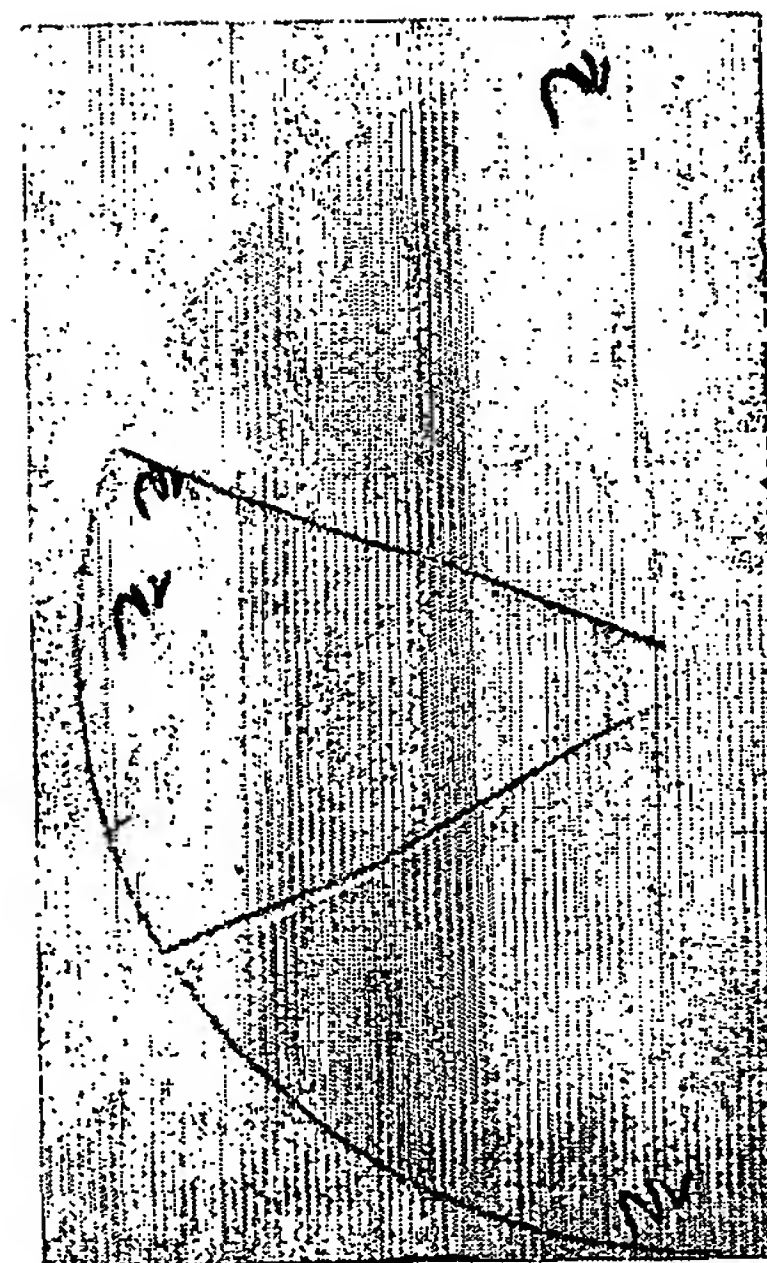
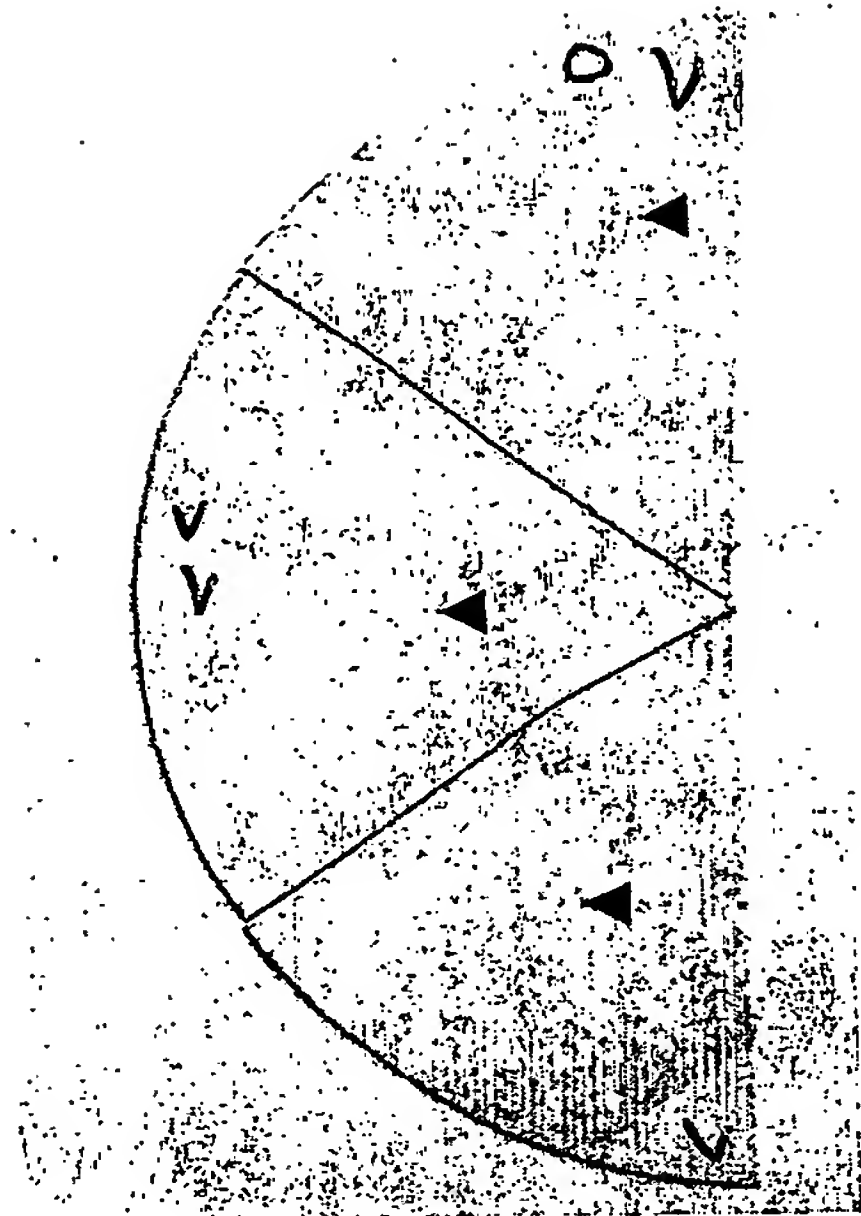


Figure 7

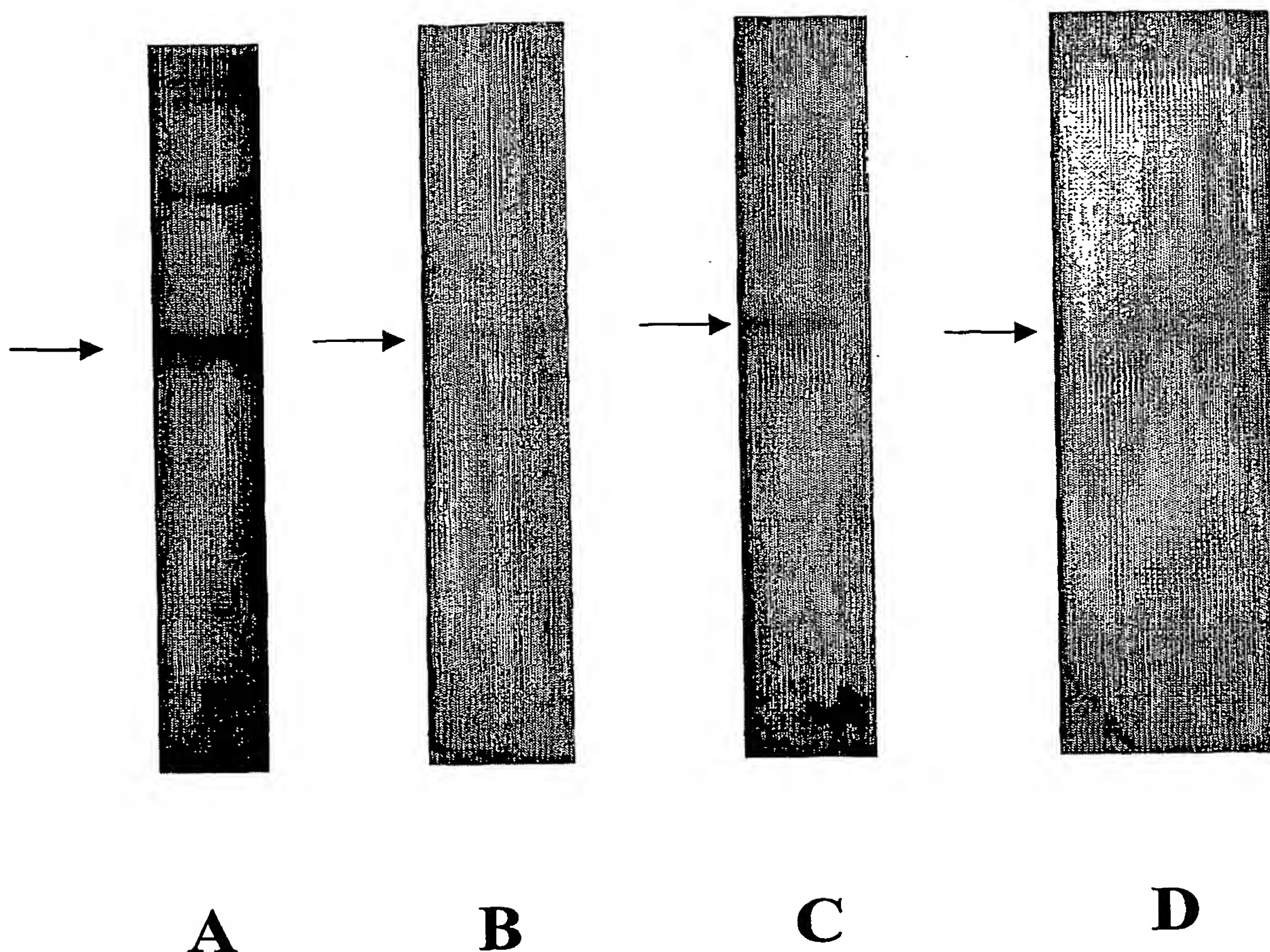


Figure 8 A

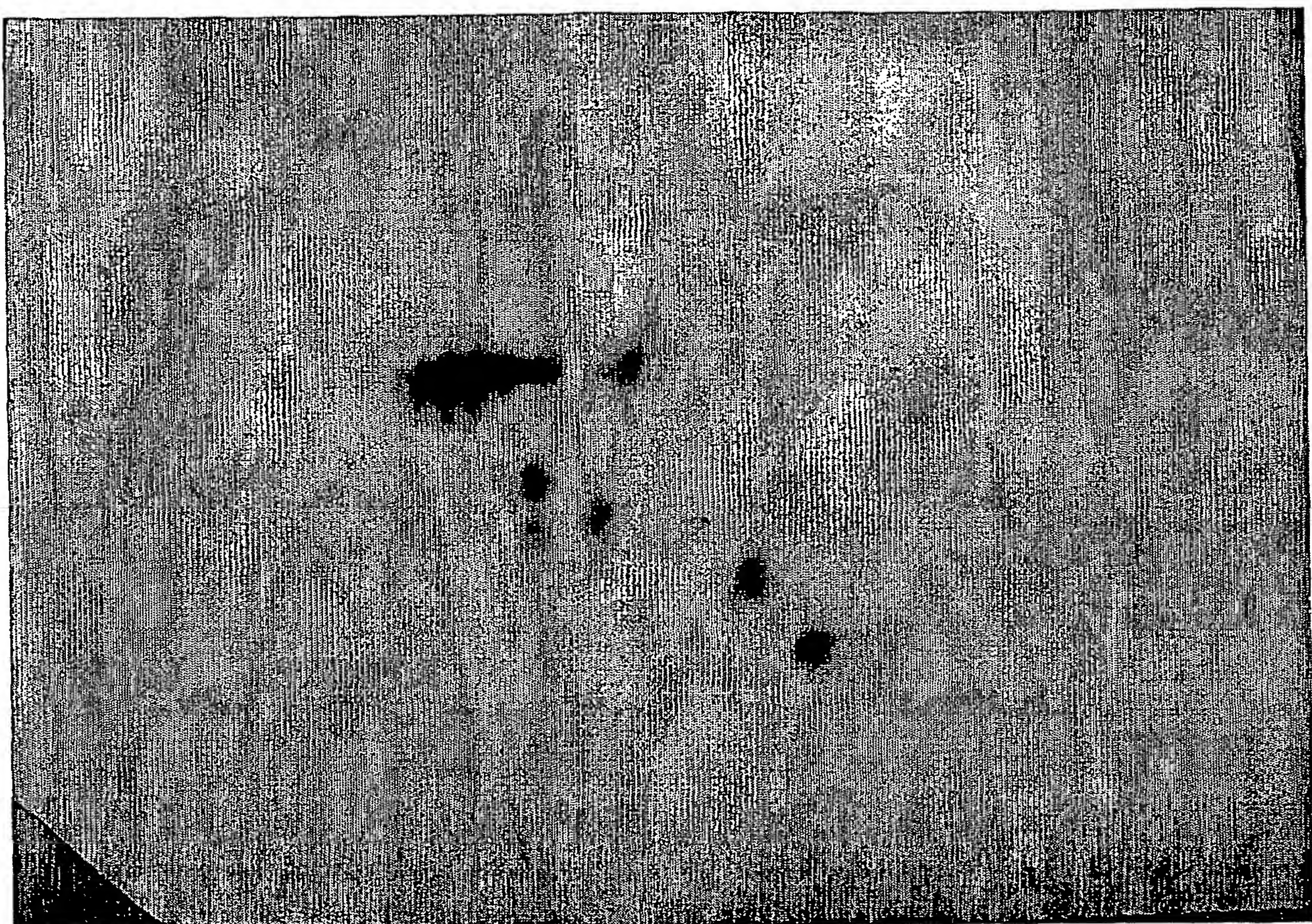


Figure 8 B

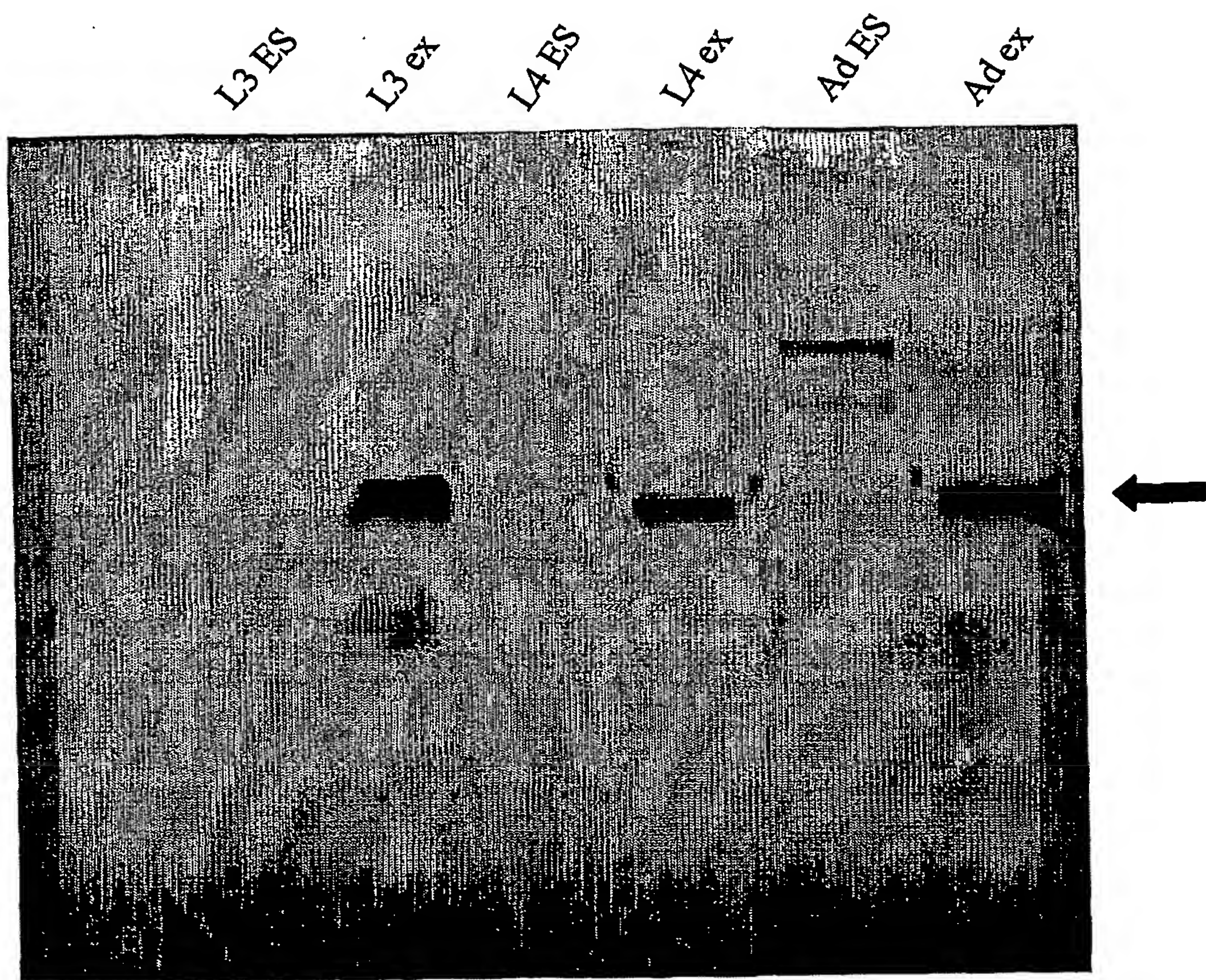


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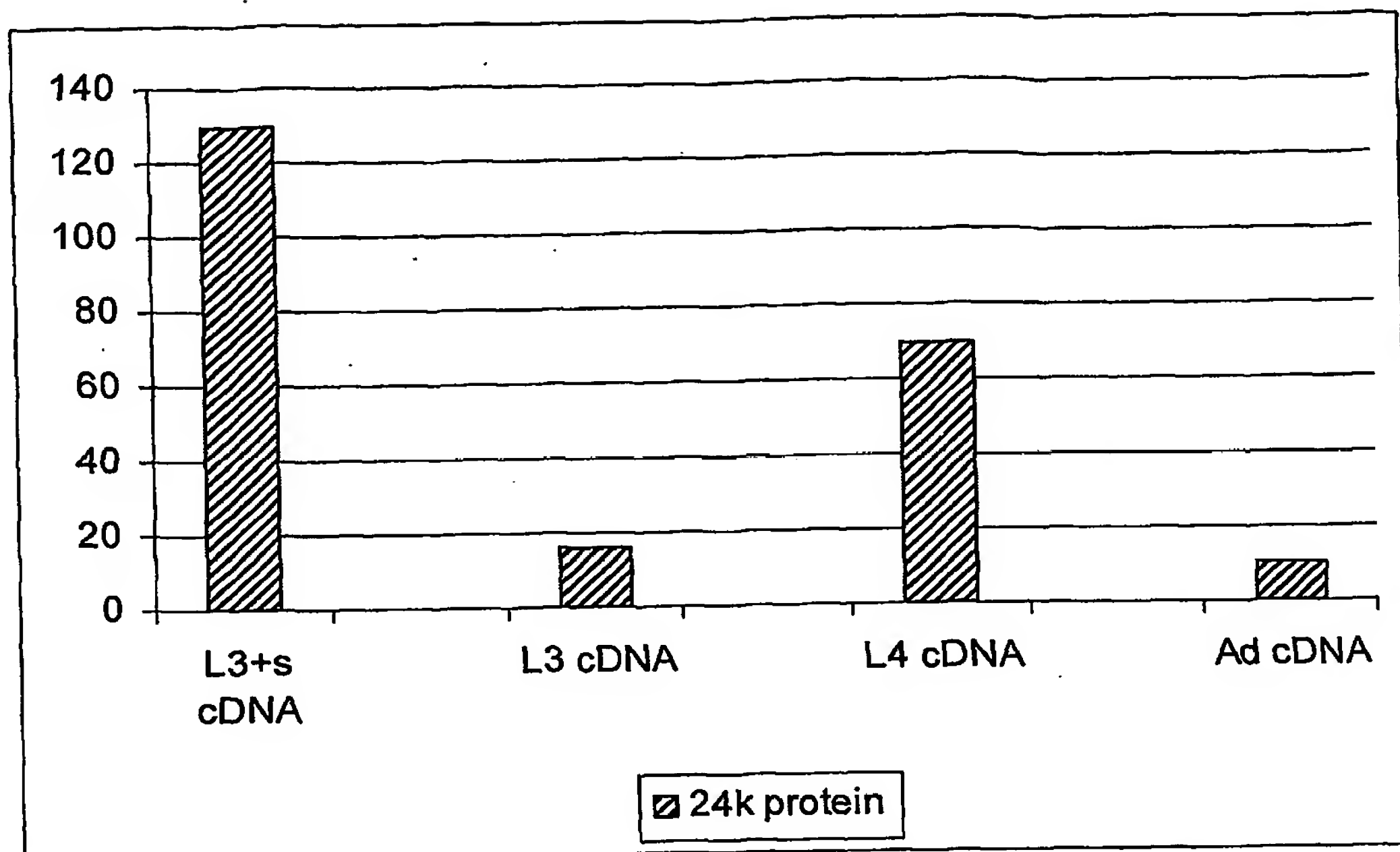


Figure 9 B

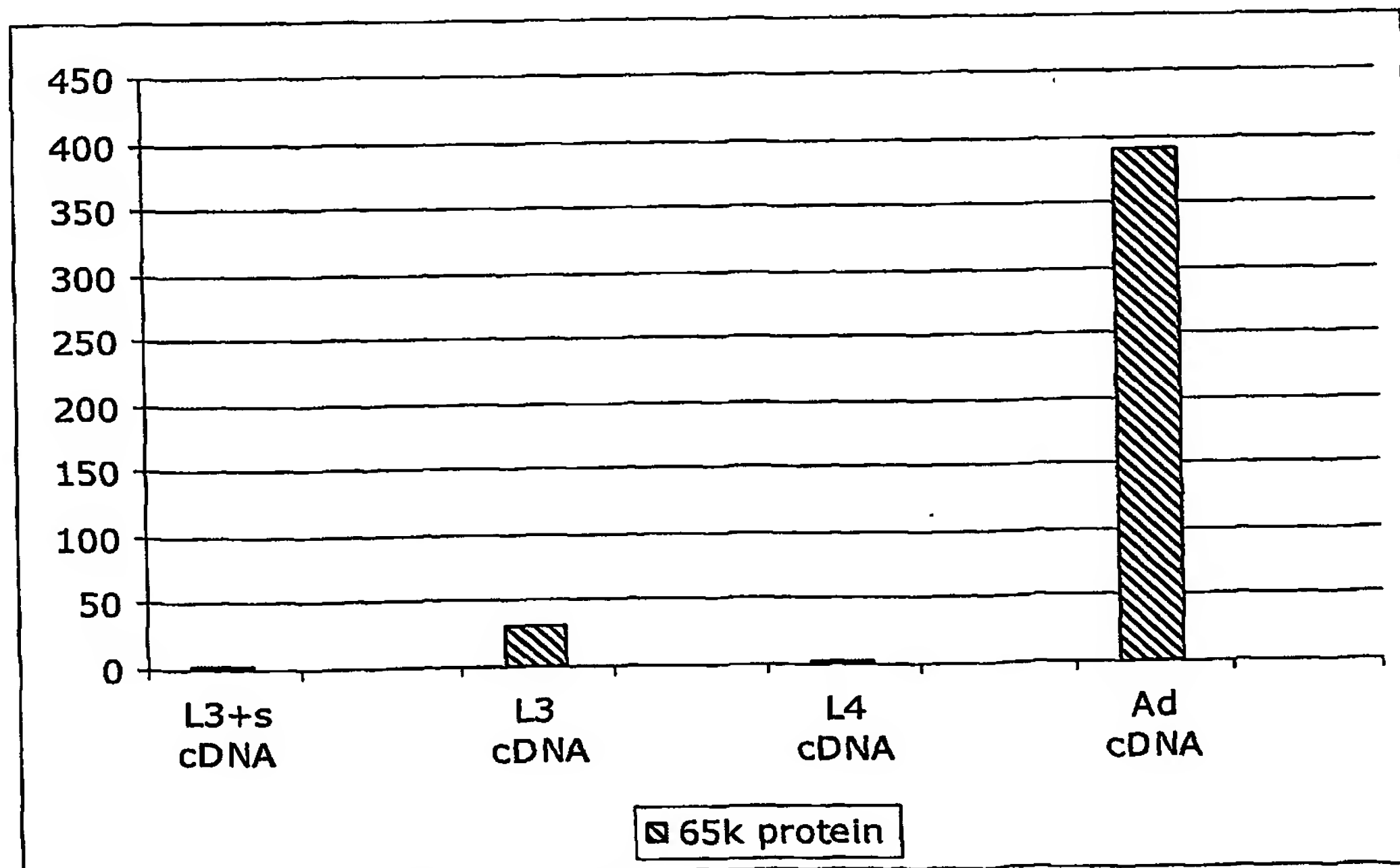
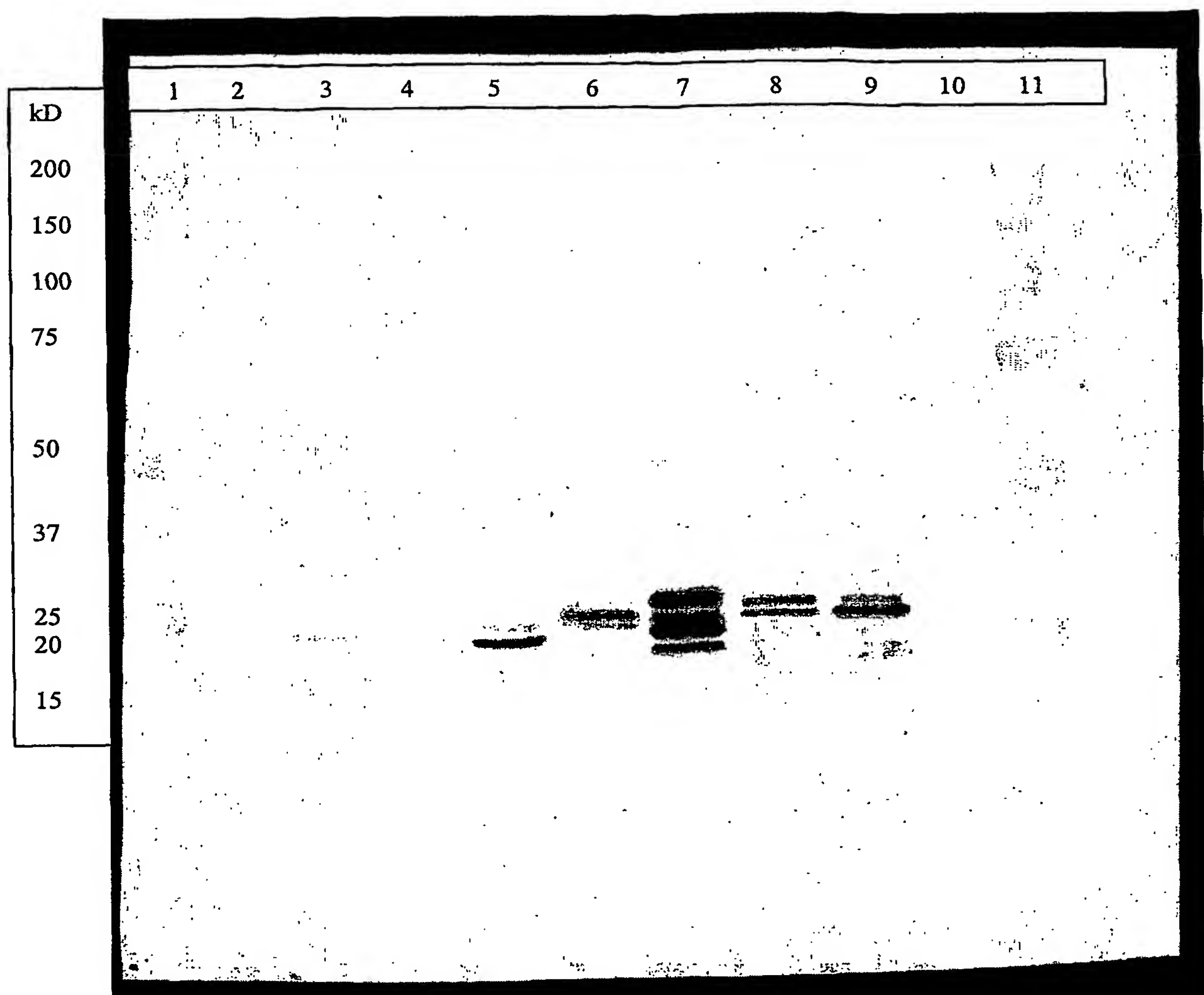


Figure 10



SEQUENCE LISTING

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 15 20 25

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Asn Gln Thr Asp Glu Ala Arg Xaa Ile Phe Leu Asp Phe His Asn Gln
30 35 40 45

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 Met Asp Trp Asp Cys Asn Leu Glu Ala Lys Ala Lys Ala Met Ile Trp
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145 150 155	
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160 165 170	
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175 180 185	
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190 195 200 205	
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210 215 220	
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35 40 45

Asp Ile Ala Gly Ala Ser Pro Leu Leu Asn Leu Thr Gly Ala Val Gln
50 55 60

Met Arg Asn Val Leu Gly Pro Ala Lys Asn Met Tyr Arg Met Asp Trp
65 70 75 80

Asp Cys Asn Leu Glu Ala Lys Ala Lys Ala Met Ile Trp Pro Cys Thr
85 90 95

Thr Pro Leu Pro Ile Asp Thr Ser Ile Pro Gln Asn Leu Ala Gln Trp
100 105 110

Leu Leu Phe Gln Asn Ser Gln Glu Xaa Glu Val Leu Thr Gln Thr Pro
115 120 125

Trp Ser Trp Val Thr Ala Ser Leu Arg Asn Leu Gln Pro Asp Thr Glu
130 135 140

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gac cgt ttc gag cgc atg ctg gaa gag ccg ttc ang cgt gtg gat cgt      95
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ttc tgc ccg atg aga gat gcg gac tgg atg agc cgt cag att atg ccc      143
  Phe Cys Pro Met Arg Asp Ala Asp Trp Met Ser Arg Gln Ile Met Pro
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nca cac ttn agg cca gaa gag ttg aag gta caa ttg gaa gtg acg      284
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Glu Ile Ile Ala Phe Ser Asp Arg Val Glu Glu Phe Lys Lys Ile Asp	
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Pro Val Leu Ala Asp Thr Asn His Gln Ile Ala Lys Asp Tyr Gly Val	
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<222> (648)..(648)
<223> n is a, c, g, or t

<220>
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<222> (662)..(662)
<223> n is a, c, g, or t

<220>
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<222> (669)..(669)
<223> n is a, c, g, or t

<220>
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<222> (682)..(682)
<223> n is a, c, g, or t

<400> 7
cta act cct tng cat cca acg cgt tgg gag ctc tnn cta tng ngg gaa
Leu Thr Pro Xaa His Pro Thr Arg Trp Glu Leu Xaa Leu Xaa Xaa Glu
1 5 10 15

48

ttg cna tgt ggt ggc gac nac tcc tgg agc ccg tca gta tcg gcg gaa
Leu Xaa Cys Gly Gly Asp Xaa Ser Trp Ser Pro Ser Val Ser Ala Glu
20 25 30

96

tcc gcg gcc gcg tcg acc gtg ggt gtg gcc ctc gcg gtc cac caa aca
Phe Ala Ala Ala Ser Thr Val Gly Val Ala Leu Ala Val His Gln Thr
35 40 45

144

ctt gac ctg ctt cct ctg aag cca cgc aag gag tac gtc ttc cgc ttt
Leu Asp Leu Leu Pro Leu Lys Pro Arg Lys Glu Tyr Val Phe Arg Phe
50 55 60

192

gaa gga nat gtt cac tcc gga atc ccg ctc cca acc gac acc acc atc
Glu Gly Xaa Val His Ser Gly Ile Pro Leu Pro Thr Asp Thr Thr Ile
65 70 75 80

240

tct cgc ata cag gct atg gta cat gtc cag atc cct gac gac cac cac
Ser Arg Ile Gln Ala Met Val His Val Gln Ile Pro Asp Asp His His
85 90 95

288

gcc att ctc aag ctg aga gat gtt cgc ttt gct act gga gaa gac gaa
Ala Ile Leu Lys Leu Arg Asp Val Arg Phe Ala Thr Gly Glu Asp Glu
100 105 110

336

cgc aga gaa ctc ttc aaa ccg atc gat gac ctg aaa atg cgc aca atc Arg Arg Glu Leu Phe Lys Pro Ile Asp Asp Leu Lys Met Arg Thr Ile 115 120 125	384
tca agg gag cac ctc gat ctc ctt gag ttg cca gtc cgt ttt gtc tac Ser Arg Glu His Leu Asp Leu Leu Glu Leu Pro Val Arg Phe Val Tyr 130 135 140	432
aag aac ggc atg att tcc gat gta atc ttt gtc gac aag gag gag acc Lys Asn Gly Met Ile Ser Asp Val Ile Phe Val Asp Lys Glu Glu Thr 145 150 155 160	480
tgg tcc cgc cag cgt gaa gcc gat ctg tca tca aca tgc tcc act tta Trp Ser Arg Gln Arg Glu Ala Asp Leu Ser Ser Thr Cys Ser Thr Leu 165 170 175	528
acc tcc aca aga tgg gac gaa ctg acn agc ttt aca atg gac agg tcc Thr Ser Thr Arg Trp Asp Glu Leu Thr Ser Phe Thr Met Asp Arg Ser 180 185 190	576
aag gtg gac ccg tng aca aac gag tac ttt cac tgg tta ccc gaa ccg Lys Val Asp Pro Xaa Thr Asn Glu Tyr Phe His Trp Leu Pro Glu Pro 195 200 205	624
aac cca ttc gaa ggg aaa ctt gtn aag gtt ggc tta cnc cgg ttn tta Asn Pro Phe Glu Gly Lys Leu Val Lys Val Gly Leu Xaa Arg Xaa Leu 210 215 220	672
aag aaa aaa nge acc ttt tgg Lys Lys Xaa Thr Phe Trp 225 230	693

<210> 8
<211> 231
<212> PRT
<213> *Ostertagia ostertagi*

<220>
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<222> (4)..(4)
<223> The 'Xaa' at location 4 stands for Trp, Ser, or Leu.

<220>
<221> misc_feature
<222> (12)..(12)
<223> The 'Xaa' at location 12 stands for Tyr, Trp, Cys, Ser, Leu, or Phe.

<220>
<221> misc_feature
<222> (14)..(14)
<223> The 'Xaa' at location 14 stands for Trp, Ser, or Leu.

<220>
<221> misc_feature
<222> (15)..(15)
<223> The 'Xaa' at location 15 stands for Arg, Gly, or Trp.

<220>
<221> misc_feature
<222> (18)..(18)
<223> The 'Xaa' at location 18 stands for Gln, Arg, Pro, or Leu.

<220>
<221> misc_feature
<222> (23)..(23)
<223> The 'Xaa' at location 23 stands for Asn, Asp, His, or Tyr.

<220>
<221> misc_feature
<222> (67)..(67)
<223> The 'Xaa' at location 67 stands for Asn, Asp, His, or Tyr.

<220>
<221> misc_feature
<222> (197)..(197)
<223> The 'Xaa' at location 197 stands for Trp, Ser, or Leu.

<220>
<221> misc_feature
<222> (221)..(221)
<223> The 'Xaa' at location 221 stands for His, Arg, Pro, or Leu.

<220>
<221> misc_feature
<222> (223)..(223)
<223> The 'Xaa' at location 223 stands for Leu, or Phe.

<220>
<221> misc_feature
<222> (228)..(228)
<223> The 'Xaa' at location 228 stands for Arg, Gly, or Trp.

<400> 8

Leu Thr Pro Xaa His Pro Thr Arg Trp Glu Leu Xaa Leu Xaa Xaa Glu
1 5 10 15

Leu Xaa Cys Gly Gly Asp Xaa Ser Trp Ser Pro Ser Val Ser Ala Glu
20 25 30

Phe Ala Ala Ala Ser Thr Val Gly Val Ala Leu Ala Val His Gln Thr
35 40 45

Leu Asp Leu Leu Pro Leu Lys Pro Arg Lys Glu Tyr Val Phe Arg Phe
50 55 60

Glu Gly Xaa Val His Ser Gly Ile Pro Leu Pro Thr Asp Thr Thr Ile
65 70 75 80

Ser Arg Ile Gln Ala Met Val His Val Gln Ile Pro Asp Asp His His
85 90 95

Ala Ile Leu Lys Leu Arg Asp Val Arg Phe Ala Thr Gly Glu Asp Glu
 100 105 110

Arg Arg Glu Leu Phe Lys Pro Ile Asp Asp Leu Lys Met Arg Thr Ile
 115 120 125

Ser Arg Glu His Leu Asp Leu Leu Glu Leu Pro Val Arg Phe Val Tyr
 130 135 140

Lys Asn Gly Met Ile Ser Asp Val Ile Phe Val Asp Lys Glu Glu Thr
 145 150 155 160

Trp Ser Arg Gln Arg Glu Ala Asp Leu Ser Ser Thr Cys Ser Thr Leu
 165 170 175

Thr Ser Thr Arg Trp Asp Glu Leu Thr Ser Phe Thr Met Asp Arg Ser
 180 185 190

Lys Val Asp Pro Xaa Thr Asn Glu Tyr Phe His Trp Leu Pro Glu Pro
 195 200 205

Asn Pro Phe Glu Gly Lys Leu Val Lys Val Gly Leu Xaa Arg Xaa Leu
 210 215 220

Lys Lys Lys Xaa Thr Phe Trp
 225 230

<210> 9
 <211> 763
 <212> DNA
 <213> *Ostertagia ostertagi*

<220>
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 <222> (11)..(706)

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 Met Ser Ala Ala Val Val Val Ala Val Leu Leu Ala Leu
 1 5 10

ttc tcc tat gcc gaa gca ggc ttt tgt tgt ccg aat agt cta agc caa
 Phe Ser Tyr Ala Glu Ala Gly Phe Cys Cys Pro Asn Ser Leu Ser Gln
 15 20 25

agt gac agc gcg agg cag att ttc ctc gat ttt cac aat gat gtt cgt
 Ser Asp Ser Ala Arg Gln Ile Phe Leu Asp Phe His Asn Asp Val Arg
 30 35 40 45

49

97

145

cga aat ata gca ctt gga aat ggt ttg ata aac tgg aca gta aat gca Arg Asn Ile Ala Leu Gly Asn Gly Leu Ile Asn Trp Thr Val Asn Ala	193
50 55 60	
gac gcg gtc att ctt ggt cca gct cag aac atg tac aaa gtg gac tgg Asp Ala Val Ile Leu Gly Pro Ala Gln Asn Met Tyr Lys Val Asp Trp	241
65 70 75	
gat tgc aac ttg gaa gaa gta gca gca caa cag att gcg cca tgc aat Asp Cys Asn Leu Glu Glu Val Ala Ala Gln Gln Ile Ala Pro Cys Asn	289
80 85 90	
gat ccc cta ccg ata aat acc agc ctg gct caa aat atc gct aga tgg Asp Pro Leu Pro Ile Asn Thr Ser Leu Ala Gln Asn Ile Ala Arg Trp	337
95 100 105	
ctg tac ttc aaa gac agt gaa gag aca gtt ctg caa caa gta tcg Leu Tyr Phe Lys Asp Ser Glu Glu Thr Val Leu Gln Gln Val Ser	385
110 115 120 125	
tgg tat tgg gtg agc gca tcg ctg gga ttt atg aaa ggc acg aaa ctt Trp Tyr Trp Val Ser Ala Ser Leu Gly Phe Met Lys Gly Thr Lys Leu	433
130 135 140	
gac caa ttt gct aac cag tgg gct gaa cct cta gca aac att gca aac Asp Gln Phe Ala Asn Gln Trp Ala Glu Pro Leu Ala Asn Ile Ala Asn	481
145 150 155	
tat aga aac cga aag gtt gga tgt gcc cat aag atc tgc ccc gct cag Tyr Arg Asn Arg Lys Val Gly Cys Ala His Lys Ile Cys Pro Ala Gln	529
160 165 170	
caa aac atg gta gta tcc tgc gtg tat gga agc ccc aaa ctt gca ccg Gln Asn Met Val Val Ser Cys Val Tyr Gly Ser Pro Lys Leu Ala Pro	577
175 180 185	
aac gaa gtt atc tgg cag gaa gga aag gct tgt gtg tgc gac gct cgt Asn Glu Val Ile Trp Gln Glu Gly Lys Ala Cys Val Cys Asp Ala Arg	625
190 195 200 205	
cca gat tca ttc tgc tgc gac aac ctg tgt gac acg cga gat gct gcg Pro Asp Ser Phe Cys Cys Asp Asn Leu Cys Asp Thr Arg Asp Ala Ala	673
210 215 220	
agt gtt cgc cac cag tgt tgc gcg tcg cca tga agcgaaaaaga aattggtagt Ser Val Arg His Gln Cys Cys Ala Ser Pro	726
225 230	
caccccgaaat aaaatattca tgcaaaaaaaaaaaaaaaa	763

<210> 10
<211> 231
<212> PRT
<213> Ostertagia ostertagi

<400> 10

Met Ser Ala Ala Val Val Val Ala Val Leu Leu Ala Leu Phe Ser Tyr
1 5 10 15

Ala Glu Ala Gly Phe Cys Cys Pro Asn Ser Leu Ser Gln Ser Asp Ser
20 25 30

Ala Arg Gln Ile Phe Leu Asp Phe His Asn Asp Val Arg Arg Asn Ile
35 40 45

Ala Leu Gly Asn Gly Leu Ile Asn Trp Thr Val Asn Ala Asp Ala Val
50 55 60

Ile Leu Gly Pro Ala Gln Asn Met Tyr Lys Val Asp Trp Asp Cys Asn
65 70 80

Leu Glu Glu Val Ala Ala Gln Ile Ala Pro Cys Asn Asp Pro Leu
85 90 95

Pro Ile Asn Thr Ser Leu Ala Gln Asn Ile Ala Arg Trp Leu Tyr Phe
100 105 110

Lys Asp Ser Glu Glu Glu Thr Val Leu Gln Gln Val Ser Trp Tyr Trp
115 120 125

Val Ser Ala Ser Leu Gly Phe Met Lys Gly Thr Lys Leu Asp Gln Phe
130 135 140

Ala Asn Gln Trp Ala Glu Pro Leu Ala Asn Ile Ala Asn Tyr Arg Asn
145 150 155 160

Arg Lys Val Gly Cys Ala His Lys Ile Cys Pro Ala Gln Gln Asn Met
165 170 175

Val Val Ser Cys Val Tyr Gly Ser Pro Lys Leu Ala Pro Asn Glu Val
180 185 190

Ile Trp Gln Glu Gly Lys Ala Cys Val Cys Asp Ala Arg Pro Asp Ser
195 200 205

Phe Cys Cys Asp Asn Leu Cys Asp Thr Arg Asp Ala Ala Ser Val Arg
210 215 220

His Gln Cys Cys Ala Ser Pro
225 230

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<210> 11
<211> 893
<212> DNA
<213> Ostertagia ostertagi

<220>
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<222> (1)..(684)

<220>
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<222> (813)..(813)
<223> n is a, c, g, or t

<220>
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<222> (858)..(858)
<223> n is a, c, g, or t

<400> 11
atg aag ttg gtc gtg ctc tgc gtt ctg tgt gga atc gct ctt gct gcc 48
Met Lys Leu Val Val Leu Cys Val Leu Cys Gly Ile Ala Leu Ala Ala
1 5 10 15

ccg aga cag aaa cgc ctt act gtg ggc acg atc gct gtc acc gga gga 96
Pro Arg Gln Lys Arg Leu Thr Val Gly Thr Ile Ala Val Thr Gly Gly
20 25 30

gtc ggc gga tcc acg ggg tgt gta gtg act gga aat gtc ctc tac gca 144
Val Gly Gly Ser Thr Gly Cys Val Val Thr Gly Asn Val Leu Tyr Ala
35 40 45

aac ggt ttc cgc ctt cgt gaa ctc aac cca tcg gag cag caa gaa ctc 192
Asn Gly Phe Arg Leu Arg Glu Leu Asn Pro Ser Glu Gln Gln Glu Leu
50 55 60

gta aac tat gag aag cag gtg gcc gac tac aaa gcg gct gtg aag caa 240
Val Asn Tyr Glu Lys Gln Val Ala Asp Tyr Lys Ala Ala Val Lys Gln
65 70 75 80

gcc ctc aag gaa cgc cag gaa agc ctg aaa tcg cgc atg gct ggt aag 288
Ala Leu Lys Glu Arg Gln Glu Ser Leu Lys Ser Arg Met Ala Gly Lys
85 90 95

aag gag aag gct gtg act ccc aag gag gaa gat cta ccc aaa gct cca 336
Lys Glu Lys Ala Val Thr Pro Lys Glu Glu Asp Leu Pro Lys Ala Pro
100 105 110

cag aag ccc tca ttc tgc act gag gac gac acc acc cag ttc tac ttt 384
Gln Lys Pro Ser Phe Cys Thr Glu Asp Asp Thr Thr Gln Phe Tyr Phe
115 120 125

gat gga tgc atg gtt cag ggc aac aag gtc tac gtt ggc aac aca ttc 432
Asp Gly Cys Met Val Gln Gly Asn Lys Val Tyr Val Gly Asn Thr Phe
130 135 140

gcg cgc gat ttg gac cag aac gag att caa gag ctg aag gag ttt gag 480
Ala Arg Asp Leu Asp Gln Asn Glu Ile Gln Glu Leu Lys Glu Phe Glu
145 150 155 160

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aag aag cag act gtc tac cag gaa tac gtc cag aag cag att caa gcg 528
 Lys Lys Gln Thr Val Tyr Gln Glu Tyr Val Gln Lys Gln Ile Gln Ala
 165 170 175

caa gtg agc aat ctg ttc ggc ggt gcc gac ttc ttt tca tcg ttc ttc 576
 Gln Val Ser Asn Leu Phe Gly Gly Ala Asp Phe Phe Ser Ser Phe Phe
 180 185 190

aac ggc gga tct gag aaa ggc tct tca acc acc act gtg gcc cca gtg 624
 Asn Gly Gly Ser Glu Lys Gly Ser Ser Thr Thr Thr Val Ala Pro Val
 195 200 205

ctt cct gaa gat gca cca gaa caa cca gct ggg ccc aac ttt tgc aca 672
 Leu Pro Glu Asp Ala Pro Glu Gln Pro Ala Gly Pro Asn Phe Cys Thr
 210 215 220

agg atc tat tga tgggtattt ttatgatgac aaagtattta aataaatgca 724
 Arg Ile Tyr
 225

gtagttgcct gttgctgtga attccacagc actcctactc acgggtgtcgta ctgggtgattt 784
 agtcacttta ttgcataat ttttatgng ttaccgcaat tcgttgtata tttgtgttat 844
 aaacattaac atcnAAAAAA aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa 893

<210> 12
<211> 227
<212> PRT
<213> Ostertagia ostertagi

<400> 12

Met Lys Leu Val Val Leu Cys Val Leu Cys Gly Ile Ala Leu Ala Ala
 1 5 10 15

Pro Arg Gln Lys Arg Leu Thr Val Gly Thr Ile Ala Val Thr Gly Gly
 20 25 30

Val Gly Gly Ser Thr Gly Cys Val Val Thr Gly Asn Val Leu Tyr Ala
 35 40 45

Asn Gly Phe Arg Leu Arg Glu Leu Asn Pro Ser Glu Gln Gln Glu Leu
 50 55 60

Val Asn Tyr Glu Lys Gln Val Ala Asp Tyr Lys Ala Ala Val Lys Gln
 65 70 75 80

Ala Leu Lys Glu Arg Gln Glu Ser Leu Lys Ser Arg Met Ala Gly Lys
 85 90 95

Lys Glu Lys Ala Val Thr Pro Lys Glu Glu Asp Leu Pro Lys Ala Pro
 100 105 110

Gln Lys Pro Ser Phe Cys Thr Glu Asp Asp Thr Thr Gln Phe Tyr Phe
 115 120 125

Asp Gly Cys Met Val Gln Gly Asn Lys Val Tyr Val Gly Asn Thr Phe
 130 135 140

Ala Arg Asp Leu Asp Gln Asn Glu Ile Gln Glu Leu Lys Glu Phe Glu
 145 150 155 160

Lys Lys Gln Thr Val Tyr Gln Glu Tyr Val Gln Lys Gln Ile Gln Ala
 165 170 175

Gln Val Ser Asn Leu Phe Gly Gly Ala Asp Phe Phe Ser Ser Phe Phe
 180 185 190

Asn Gly Gly Ser Glu Lys Gly Ser Ser Thr Thr Thr Val Ala Pro Val
 195 200 205

Leu Pro Glu Asp Ala Pro Glu Gln Pro Ala Gly Pro Asn Phe Cys Thr
 210 215 220

Arg Ile Tyr
 225

<210> 13
<211> 1761
<212> DNA
<213> Ostertagia ostertagi

<220>
<221> CDS
<222> (1)..(1725)

<400> 13
atg agg ctg ata ttg ctc att tta ctc ttg gtt gtt gcc act aat ggg 48
Met Arg Leu Ile Leu Leu Ile Leu Leu Val Val Ala Thr Asn Gly
1 5 10 15

ggc ata att gac aaa ctg aaa gga ttg ttc act gga gaa ggc ggc ttt 96
Gly Ile Ile Asp Lys Leu Lys Gly Leu Phe Thr Gly Glu Gly Gly Phe
20 25 30

gga caa aaa gtg aag aat gca act gct gtt ggc ttc aaa aag ctc ttc 144
Gly Gln Lys Val Lys Asn Ala Thr Ala Val Gly Phe Lys Lys Leu Phe
35 40 45

gaa aac acg gca ctc ttc aga atc aat gat aag atc agg agc atg aag	192
Glu Asn Thr Ala Leu Phe Arg Ile Asn Asp Lys Ile Arg Ser Met Lys	
50 55 60	
gaa aaa gtg ttg aag acc ttg gaa cta tca cca gca atg atg aag tca	240
Glu Lys Val Leu Lys Thr Leu Glu Leu Ser Pro Ala Met Met Lys Ser	
65 70 75 80	
ctg caa kmg agg cta rwg aaw tsg cgr cck yct rma grw cga yma wrt	288
Leu Gln Xaa Arg Leu Xaa Xaa Arg Xaa Xaa Xaa Xaa Arg Xaa Xaa	
85 90 95	
rsr mga gmt sss aga crc gtw kka ygc rag gtc art aaa aat agt gag	336
Xaa Xaa Xaa Arg Xaa Xaa Xaa Val Xaa Lys Asn Ser Glu	
100 105 110	
gtt gac caa tac ctc tac caa ggc gac atg gtt tta aca gag gag caa	384
Val Asp Gln Tyr Leu Tyr Gln Gly Asp Met Val Leu Thr Glu Glu Gln	
115 120 125	
gcc gat gag atc gtt gag gac ata gaa gat cag gtc gcc ggt gga aat	432
Ala Asp Glu Ile Val Glu Asp Ile Glu Asp Gln Val Ala Gly Gly Asn	
130 135 140	
cgt aca aaa cgt caa gca ttc aag gat cat aaa tat ccc aaa acg ttg	480
Arg Thr Lys Arg Gln Ala Phe Lys Asp His Lys Tyr Pro Lys Thr Leu	
145 150 155 160	
tgg tca caa gga gtc aac tac tac ttc cat gat atg gcc agt aag cag	528
Trp Ser Gln Gly Val Asn Tyr Tyr Phe His Asp Met Ala Ser Lys Gln	
165 170 175	
atg aaa agc gta ttc gta aaa gga gcg aaa tgg tgg gaa aag gac acg	576
Met Lys Ser Val Phe Val Lys Gly Ala Lys Trp Trp Glu Lys Asp Thr	
180 185 190	
tgt atc aat ttc acg gag aac cgt tct gcc gaa gac cga att atg gta	624
Cys Ile Asn Phe Thr Glu Asn Arg Ser Ala Glu Asp Arg Ile Met Val	
195 200 205	
ttc cca cag aaa gga tgt tgg tca aat ata gga aaa atc ggt ggc gaa	672
Phe Pro Gln Lys Gly Cys Trp Ser Asn Ile Gly Lys Ile Gly Gly Glu	
210 215 220	
caa aag att tcg ttg gga gga ggt tgt cat tcg gta tcc att gct gcg	720
Gln Lys Ile Ser Leu Gly Gly Gly Cys His Ser Val Ser Ile Ala Ala	
225 230 235 240	
cat gag atc ggt cac gca att gga ttc ttc cat act atg tcc cgt cac	768
His Glu Ile Gly His Ala Ile Gly Phe Phe His Thr Met Ser Arg His	
245 250 255	
gat cgc gat gaa ttc atc acc gta aac atg cac aat gtt gat gta cac	816
Asp Arg Asp Glu Phe Ile Thr Val Asn Met His Asn Val Asp Val His	
260 265 270	
tgg ctg agt caa ttt aat aaa gaa acg acg aag aga aat gat aat tat	864
Trp Leu Ser Gln Phe Asn Lys Glu Thr Thr Lys Arg Asn Asp Asn Tyr	
275 280 285	

gga atg acg tac gac tac ggt agc att atg cat tac ggt gga acc agt Gly Met Thr Tyr Asp Tyr Gly Ser Ile Met His Tyr Gly Gly Thr Ser 290 295 300	912
gca tcg tac aat aat aag cca aca atg gtg ccg ttt gat gtg gac tat Ala Ser Tyr Asn Asn Lys Pro Thr Met Val Pro Phe Asp Val Asp Tyr 305 310 315 320	960
cag caa acc ctt ggc tct cca ttc att tct ttc att gaa ctt tcc atg Gln Gln Thr Leu Gly Ser Pro Phe Ile Ser Phe Ile Glu Leu Ser Met 325 330 335	1008
att aat gaa cac tac aaa tgc aaa gag aac tgc aat cca gct aag tcg Ile Asn Glu His Tyr Lys Cys Lys Glu Asn Cys Asn Pro Ala Lys Ser 340 345 350	1056
gct aaa tgc gaa atg ggc gga ttc cct cat ccc cga gac tgc agc aaa Ala Lys Cys Glu Met Gly Phe Pro His Pro Arg Asp Cys Ser Lys 355 360 365	1104
tgt atc tgt cct ggt gga tac gcc gga gct cga tgc acc gaa aga cca Cys Ile Cys Pro Gly Gly Tyr Ala Gly Ala Arg Cys Thr Glu Arg Pro 370 375 380	1152
tca ggg tgt ggc agt gca att caa gct tcg tcc gat tgg aag acc tta Ser Gly Cys Gly Ser Ala Ile Gln Ala Ser Ser Asp Trp Lys Thr Leu 385 390 395 400	1200
caa gat acc ctt ggc aag gat gat gat gaa gaa cga gag gac ttc gag Gln Asp Thr Leu Gly Lys Asp Asp Asp Glu Glu Arg Glu Asp Phe Glu 405 410 415	1248
aca tgt aat tac tgg att gaa tct cct gcc gga acm gaa atc gaa gtc Thr Cys Asn Tyr Trp Ile Glu Ser Pro Ala Gly Xaa Glu Ile Glu Val 420 425 430	1296
agg tta ttg gat ttc acg agg ggt gtt tct gtc gat gga tgc aaa ttt Arg Leu Leu Asp Phe Thr Arg Gly Val Ser Val Asp Gly Cys Lys Phe 435 440 445	1344
gcc ggt gta gag atc aag acc aat aag gat caa aca ctc act ggc tac Ala Gly Val Glu Ile Lys Thr Asn Lys Asp Gln Thr Leu Thr Gly Tyr 450 455 460	1392
agg ttc tgc aca gct ggc gca gct ggc ata gca ctt cgt tct tac acg Arg Phe Cys Thr Ala Gly Ala Ala Gly Ile Ala Leu Arg Ser Tyr Thr 465 470 475 480	1440
aat cgc gtc cca ata atg aca tac aac aga ttt ggt caa tcg acg act Asn Arg Val Pro Ile Met Thr Tyr Asn Arg Phe Gly Gln Ser Thr Thr 485 490 495	1488
gtt ctc gaa tac cga cac gtt ccg gcg agt gcg cca aga acg ccc tca Val Leu Glu Tyr Arg His Val Pro Ala Ser Ala Pro Arg Thr Pro Ser 500 505 510	1536
cct cca tct gct aca act cgt gct tct att act act act act acg Pro Pro Ser Ala Thr Thr Arg Ala Ser Ile Thr Thr Thr Thr Thr 515 520 525	1584

aag aaa ccc agc tct act gct gcc ttt aaa tgc gag gac aac cac act	1632
Lys Lys Pro Ser Ser Thr Ala Ala Phe Lys Cys Glu Asp Asn His Thr	
530 535 540	
tgt ccc tca ctt gta gcg agc ggt ttc tgc aaa ggg cca ctc tca gag	1680
Cys Pro Ser Leu Val Ala Ser Gly Phe Cys Lys Gly Pro Leu Ser Glu	
545 550 555 560	
gct acc aag aag aaa gtg tgt cca aag tcg tgt gga ctc tgc tga	1725
Ala Thr Lys Lys Val Cys Pro Lys Ser Cys Gly Leu Cys	
565 570	
tacaacactt tctctgtaat aaaatctgaa caattc	1761

- <210> 14
- <211> 574
- <212> PRT
- <213> Ostertagia ostertagi

- <220>
- <221> misc_feature
- <222> (83)..(83)
- <223> The 'Xaa' at location 83 stands for Glu, Ala, or Ser.

- <220>
- <221> misc_feature
- <222> (86)..(86)
- <223> The 'Xaa' at location 86 stands for Glu, Val, Lys, or Met.

- <220>
- <221> misc_feature
- <222> (87)..(87)
- <223> The 'Xaa' at location 87 stands for Lys, or Asn.

- <220>
- <221> misc_feature
- <222> (88)..(88)
- <223> The 'Xaa' at location 88 stands for Trp, or Ser.

- <220>
- <221> misc_feature
- <222> (90)..(90)
- <223> The 'Xaa' at location 90 stands for Pro.

- <220>
- <221> misc_feature
- <222> (91)..(91)
- <223> The 'Xaa' at location 91 stands for Pro, or Ser.

- <220>
- <221> misc_feature
- <222> (92)..(92)
- <223> The 'Xaa' at location 92 stands for Glu, Ala, Lys, or Thr.

- <220>
- <221> misc_feature
- <222> (93)..(93)
- <223> The 'Xaa' at location 93 stands for Gly, Glu, or Asp.

<220>
<221> misc_feature
<222> (95)..(95)
<223> The 'Xaa' at location 95 stands for Gln, Pro, or Ser.

<220>
<221> misc_feature
<222> (96)..(96)
<223> The 'Xaa' at location 96 stands for Ser, Asn, Cys, or Tyr.

<220>
<221> misc_feature
<222> (97)..(97)
<223> The 'Xaa' at location 97 stands for Gly, Ala, Arg, or Thr.

<220>
<221> misc_feature
<222> (98)..(98)
<223> The 'Xaa' at location 98 stands for Arg.

<220>
<221> misc_feature
<222> (99)..(99)
<223> The 'Xaa' at location 99 stands for Asp, or Ala.

<220>
<221> misc_feature
<222> (100)..(100)
<223> The 'Xaa' at location 100 stands for Gly, Ala, Arg, or Pro.

<220>
<221> misc_feature
<222> (102)..(102)
<223> The 'Xaa' at location 102 stands for Arg, or His.

<220>
<221> misc_feature
<222> (103)..(103)
<223> The 'Xaa' at location 103 stands for Val.

<220>
<221> misc_feature
<222> (104)..(104)
<223> The 'Xaa' at location 104 stands for Gly, Val, or Leu.

<220>
<221> misc_feature
<222> (105)..(105)
<223> The 'Xaa' at location 105 stands for Arg, or Cys.

<220>
<221> misc_feature
<222> (106)..(106)
<223> The 'Xaa' at location 106 stands for Glu, or Lys.

<220>
<221> misc_feature
<222> (108)..(108)
<223> The 'Xaa' at location 108 stands for Ser, or Asn.

<220>

<221> misc_feature

<222> (428)...(428)

<223> The 'Xaa' at location 428 stands for Thr.

<400> 14

Met Arg Leu Ile Leu Leu Ile Leu Leu Val Val Ala Thr Asn Gly
1 5 10 15

Gly Ile Ile Asp Lys Leu Lys Gly Leu Phe Thr Gly Glu Gly Gly Phe
20 25 30

Gly Gln Lys Val Lys Asn Ala Thr Ala Val Gly Phe Lys Lys Leu Phe
35 40 45

Glu Asn Thr Ala Leu Phe Arg Ile Asn Asp Lys Ile Arg Ser Met Lys
50 55 60

Glu Lys Val Leu Lys Thr Leu Glu Leu Ser Pro Ala Met Met Lys Ser
65 70 75 80

Leu Gln Xaa Arg Leu Xaa Xaa Xaa Arg Xaa Xaa Xaa Xaa Arg Xaa Xaa
85 90 95

Xaa Xaa Xaa Xaa Arg Xaa Xaa Xaa Xaa Val Xaa Lys Asn Ser Glu
100 105 110

Val Asp Gln Tyr Leu Tyr Gln Gly Asp Met Val Leu Thr Glu Glu Gln
115 120 125

Ala Asp Glu Ile Val Glu Asp Ile Glu Asp Gln Val Ala Gly Gly Asn
130 135 140

Arg Thr Lys Arg Gln Ala Phe Lys Asp His Lys Tyr Pro Lys Thr Leu
145 150 155 160

Trp Ser Gln Gly Val Asn Tyr Tyr Phe His Asp Met Ala Ser Lys Gln
165 170 175

Met Lys Ser Val Phe Val Lys Gly Ala Lys Trp Trp Glu Lys Asp Thr
180 185 190

Cys Ile Asn Phe Thr Glu Asn Arg Ser Ala Glu Asp Arg Ile Met Val
195 200 205

Phe Pro Gln Lys Gly Cys Trp Ser Asn Ile Gly Lys Ile Gly Gly Glu
210 215 220

Gln Lys Ile Ser Leu Gly Gly Cys His Ser Val Ser Ile Ala Ala
225 230 235 240

His Glu Ile Gly His Ala Ile Gly Phe Phe His Thr Met Ser Arg His
245 250 255

Asp Arg Asp Glu Phe Ile Thr Val Asn Met His Asn Val Asp Val His
260 265 270

Trp Leu Ser Gln Phe Asn Lys Glu Thr Thr Lys Arg Asn Asp Asn Tyr
275 280 285

Gly Met Thr Tyr Asp Tyr Gly Ser Ile Met His Tyr Gly Gly Thr Ser
290 295 300

Ala Ser Tyr Asn Asn Lys Pro Thr Met Val Pro Phe Asp Val Asp Tyr
305 310 315 320

Gln Gln Thr Leu Gly Ser Pro Phe Ile Ser Phe Ile Glu Leu Ser Met
325 330 335

Ile Asn Glu His Tyr Lys Cys Lys Glu Asn Cys Asn Pro Ala Lys Ser
340 345 350

Ala Lys Cys Glu Met Gly Gly Phe Pro His Pro Arg Asp Cys Ser Lys
355 360 365

Cys Ile Cys Pro Gly Gly Tyr Ala Gly Ala Arg Cys Thr Glu Arg Pro
370 375 380

Ser Gly Cys Gly Ser Ala Ile Gln Ala Ser Ser Asp Trp Lys Thr Leu
385 390 395 400

Gln Asp Thr Leu Gly Lys Asp Asp Asp Glu Glu Arg Glu Asp Phe Glu
405 410 415

Thr Cys Asn Tyr Trp Ile Glu Ser Pro Ala Gly Xaa Glu Ile Glu Val
420 425 430

Arg Leu Leu Asp Phe Thr Arg Gly Val Ser Val Asp Gly Cys Lys Phe
435 440 445

Ala Gly Val Glu Ile Lys Thr Asn Lys Asp Gln Thr Leu Thr Gly Tyr
 450 455 460

Arg Phe Cys Thr Ala Gly Ala Ala Gly Ile Ala Leu Arg Ser Tyr Thr
 465 470 475 480

Asn Arg Val Pro Ile Met Thr Tyr Asn Arg Phe Gly Gln Ser Thr Thr
 485 490 495

Val Leu Glu Tyr Arg His Val Pro Ala Ser Ala Pro Arg Thr Pro Ser
 500 505 510

Pro Pro Ser Ala Thr Thr Arg Ala Ser Ile Thr Thr Thr Thr Thr Thr
 515 520 525

Lys Lys Pro Ser Ser Thr Ala Ala Phe Lys Cys Glu Asp Asn His Thr
 530 535 540

Cys Pro Ser Leu Val Ala Ser Gly Phe Cys Lys Gly Pro Leu Ser Glu
 545 550 555 560

Ala Thr Lys Lys Lys Val Cys Pro Lys Ser Cys Gly Leu Cys
 565 570

<210> 15

<211> 24

<212> DNA

<213> Artificial

<220>

<223> primer: Lambdagt11F

<220>

<221> misc_feature

<223> Lambdagt11F

<400> 15

ggtgtggcgacg actccctggag cccg

24

<210> 16

<211> 24

<212> DNA

<213> Artificial

<220>

<223> primer: Lambdagt11R

<400> 16

tttgacaccagg accaactgggt aatg

24

<210> 17
<211> 20
<212> DNA
<213> Artificial

<220>
<223> primer: SP6

<400> 17
atttaggtga cactatagaa

20

<210> 18
<211> 22
<212> DNA
<213> Artificial

<220>
<223> primer: T7

<400> 18
gtaatacgac tcactatagg gc

22

<210> 19
<211> 21
<212> DNA
<213> Artificial

<220>
<223> primer: 24kForw

<400> 19
gaattcatga agttggtcgt g

21

<210> 20
<211> 22
<212> DNA
<213> Artificial

<220>
<223> primer: 24kRev

<400> 20
ctcgagtc当地 tagatccttg tg

22

<210> 21
<211> 36
<212> DNA
<213> Artificial

<220>
<223> primer: AAP

<220>
<221> misc_feature
<222> (24)..(25)
<223> n = Inosine

<220>
<221> misc_feature
<222> (29)..(30)
<223> n = Inosine

<220>
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<222> (34)..(35)
<223> n = Inosine

<400> 21
ggccacgcgt cgactagtagc gggnnnnnnn gggnnng

36

<210> 22
<211> 32
<212> DNA
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<220>
<223> primer: UAP

<400> 22
cuacuacuac uaggccacgc gtcgactagt ac

32

<210> 23
<211> 21
<212> DNA
<213> Artificial

<220>
<223> primer: 65Rev1

<400> 23
cagcaatgga taccgaatga c

21

<210> 24
<211> 22
<212> DNA
<213> Artificial

<220>
<223> primer: 65Rev2

<400> 24
agtgacttca tcattgctgg tg

22

<210> 25
<211> 21
<212> DNA
<213> Artificial

<220>
<223> primer: 65kForw

<400> 25
tgatgatgaa gaacgagagg a

21

<210> 26
<211> 30
<212> DNA
<213> Artificial

<220>
<223> primer: For65

<400> 26
ggatccatga ggctgatatt gtcattttta

30

<210> 27
<211> 27
<212> DNA
<213> Artificial

<220>
<223> primer: Rev65

<400> 27
ctcgaggcag agtccacacg actttgg

27